Appropriateness of clinical severity classification of new WHO childhood pneumonia guidance: a multi-hospital, retrospective, cohort study

Ambrose Agweyu, Richard J Lilford, Mike English, for the Clinical Information Network Author Group*

Summary

Background Management of pneumonia in many low-income and middle-income countries is based on WHO guidelines that classify children according to clinical signs that define thresholds of risk. We aimed to establish whether some children categorised as eligible for outpatient treatment might have a risk of death warranting their treatment in hospital.

Methods We did a retrospective cohort study of children aged 2–59 months admitted to one of 14 hospitals in Kenya with pneumonia between March 1, 2014, and Feb 29, 2016, before revised WHO pneumonia guidelines were adopted in the country. We modelled associations with inpatient mortality using logistic regression and calculated absolute risks of mortality for presenting clinical features among children who would, as part of revised WHO pneumonia guidelines, be eligible for outpatient treatment (non-severe pneumonia).

Findings We assessed 16,162 children who were admitted to hospital in this period. 832 (5%) of 16,031 children died. Among groups defined according to new WHO guidelines, 321 (3%) of 11,788 patients with non-severe pneumonia died compared with 488 (14%) of 3,434 patients with severe pneumonia. Three characteristics were strongly associated with death of children retrospectively classified as having non-severe pneumonia: severe pallor (adjusted risk ratio 5·9, 95% CI 5·1–6·8), mild to moderate pallor (3·4, 3·0–3·8), and weight-for-age Z score (WAZ) less than –3 SD (3·8, 3·4–4·3). Additional factors that were independently associated with death were: WAZ less than –2 to –3 SD, age younger than 12 months, lower chest wall indrawing, respiratory rate of 70 breaths per min or more, female sex, admission to hospital in a malaria endemic region, moderate dehydration, and an axillary temperature of 39°C or more.

Interpretation In settings of high mortality, WAZ less than –3 SD or any degree of pallor among children with non-severe pneumonia was associated with a clinically important risk of death. Our data suggest that admission to hospital should not be denied to children with these signs and we urge clinicians to consider these risk factors in addition to WHO criteria in their decision making.

Funding Wellcome Trust.

Copyright © The Author(s). Published by Elsevier Ltd. This is an Open Access article under the CC BY 4.0 license.

Introduction

Management of children with pneumonia is based on their anticipated risk of poor outcome. In many low-income and middle-income countries that use WHO case management guidelines, hospital admission is recommended when a child crosses the threshold from a non-severe to a severe pneumonia classification according to revised WHO 2013 definitions (table 1).1 Hospital care should allow for prompt identification of signs of clinical deterioration and timely intervention with appropriate investigations, treatment, and supportive care including oxygen, fluids, and feeds. Children admitted to hospital could also benefit from expert review that can detect other causes of illness such as heart disease, which might be misdiagnosed as pneumonia by junior clinicians. WHO’s revised classification of pneumonia included a change from three to two severity strata, such that children with lower chest wall indrawing are now classed as having non-severe pneumonia, whereas a child with this sign would have automatically been assigned to the severe category under the previous set of criteria.1 Other events that have affected the classification of pneumonia include the widespread availability of conjugate vaccines against Streptococcus pneumoniae and Haemophilus influenzae type b (Hib);2 declining prevalence of vertically acquired HIV;3 and a technical update of WHO guidelines for the management of severe acute malnutrition in infants and children, which advises the use of weight-for-height Z score (WHZ) and mid-upper-arm circumference to replace weight-for-age Z score (WAZ) for the diagnosis of severe malnutrition requiring inpatient management.4 Although studies suggested that the previous WHO clinical algorithm performed similarly across different geographical locations,5 concerns have been raised regarding the applicability of new WHO pneumonia guidelines in high mortality settings.6,7 In this study, we assess risk factors for death in children treated in hospital with pneumonia in Kenya before
Implementation of the revised WHO pneumonia guidelines. This analysis allows us to explore outcomes in a population of children for whom admission to hospital would no longer be recommended and investigate additional clinical features that might improve risk assessment.

Methods

Study design and participants

We did a retrospective cohort study using data from an established network of 14 purposely selected public hospitals situated in regions of high and low malaria transmission in Kenya.5 Inpatient records for all children aged 2–59 months admitted to hospital with pneumonia between March 1, 2014, and Feb 29, 2016, were included. We excluded children with documented comorbidities because these are specifically excluded from the WHO pneumonia case management algorithm. Comorbidities include suspected or confirmed meningitis, HIV exposure or infection, severe acute malnutrition (typically identified by presence of visible severe wasting or oedema, skin changes of kwashiorkor, and mid-upper-arm circumference less than 11.5 cm or WHZ less than −3 SD from WHO reference charts), and chronic cardiopulmonary illnesses. Thus, we studied children in whom a clinical diagnosis of pneumonia was the primary indication for antibiotic therapy. WHO uses the terms pneumonia and severe pneumonia to denote groups requiring different treatments. To avoid confusion, we use the term pneumonia to refer to all children with a clinical diagnosis of pneumonia and the terms non-severe and severe pneumonia to refer to children who are recommended outpatient and inpatient treatment respectively by WHO (table 1).

The participating hospitals are typical of district-level health facilities in Kenya and in sub-Saharan Africa, where diagnostic capacity is low. Pulse oximetry data were not available for over 50% of patients and in some hospitals data were not available at all. In line with Kenyan policy recommendations, patients with uncomplicated pneumonia do not routinely have x-rays. In the few cases in which x-rays are done, no structured tools exist for data capture. The Hib and pneumococcal conjugate vaccines were introduced to the national routine immunisation schedule in 2001 and January, 2011, respectively. Survey data report over 80% coverage for three doses of these vaccines among children aged 12 months.6 We restricted our analysis to children who were born after the introduction of Hib and pneumococcal conjugate vaccines.

Data were extracted from patient records devoid of names and contact details that could identify individuals whose information was collected. Data entry was done following patient discharge from hospital or death; thus, the exercise did not interfere with routine patient care or pose apparent additional risk to patients. Ethics approval for the primary study was obtained from the Kenya Medical Research Institute (KEMRI) National Ethical Review Committee. The Ministry of Health and participating hospitals gave permission for the study.

Evidence before this study

We searched MEDLINE for relevant articles published until Jan 16, 2017, on risk factors for community-acquired pneumonia in children in low-income and middle-income countries by use of the terms “respiratory tract infection” [MeSH terms] OR “pneumonia” AND “(child OR paediatric OR pediatric)” AND (“mortality OR death”). The search was not restricted by date or language. We retrieved a systematic review of 77 observational studies (198 359 children) on risk factors for death from acute lower respiratory infections in children in low-income and middle-income countries.

Added value of this study

Published evidence on risk factors for community-acquired childhood pneumonia comes from studies almost entirely done in the pre-pneumococcal vaccine period. The clinical cause of childhood pneumonia is believed to be changing with increasingly high coverage of conjugate vaccines targeting previously dominant causes of pneumonia. This study—the largest published individual analysis of risk factors for mortality among children admitted to hospital with pneumonia—presents findings from a high mortality setting with a high prevalence of comorbidity. We also present unique data on risk of death for subpopulations of children classified as having non-severe pneumonia, who, under existing guidance, might have a level of risk warranting inpatient care.

Implications of all the available evidence

The risk factors for pneumonia mortality include severe pneumonia (previously very severe pneumonia), low weight-for-age Z score (WAZ), female sex, age younger than 12 months, Pneumocystis carinii infection, HIV infection, young maternal age, low maternal education, low socioeconomic status, secondhand cigarette smoke exposure, indoor air pollution, childhood immunisation status not up to date, non-attendance of antenatal care, and from our study, mild and moderate pallor, respiratory rate of 70 breaths per min or more, admission to hospital in a malaria endemic region, moderate dehydration, and an axillary temperature of 39°C or above. The risk of death for children classified as having non-severe pneumonia with mild to moderate pallor, severe pallor, or low WAZ is similar to that observed for children with WHO-defined severe pneumonia. In high mortality settings, a need exists to re-examine criteria for admission, and clinicians should treat the WHO severe category as but one factor to consider in clinical decision making.
Procedures
Methods of collection and cleaning of data in the clinical information network are reported in detail elsewhere. Clinical data for children admitted to hospitals within the network are captured through structured Paediatric Admission Record forms, approved by the Ministry of Health, that prompt the clinician with a checklist of fields including patient biodata, information gathered during clinical assessment, admission and discharge diagnoses, treatments, and outcome (survival or death). The network supports one data clerk in each hospital to extract data from medical records, nursing charts, treatment charts, and available laboratory reports each day as children are discharged. Data are abstracted from inpatient records into the primary data collection tool developed in Research Electronic Data Capture (REDCap) with error checks at point of entry by automated daily review, and with regular external data quality assurance.

Statistical analysis
Categorical data were tabulated and summarised as proportions, whereas continuous variables were reported as mean (SD) or median (IQR) as appropriate. Univariate associations with mortality were calculated for demographic and clinical characteristics with at least 50% of data available. These variables were subsequently considered for inclusion in multivariable logistic regression models restricted to complete cases. Patients’ age and sex were included in multivariable models a priori. Hospital identity was included in models as a random effect, whereas location was grouped as a binary variable on the basis of high versus low prevalence of malaria and included as a fixed effect. Other patient characteristics associated with mortality and a p value of less than 0·2 in univariate analyses were added sequentially, beginning with characteristics with the strongest association from univariate models. Likelihood ratio tests were done on addition of each variable to identify the variables to be retained (Model 1). Subsequently, multiple imputation by chained equations (generating ten imputed datasets) was done under missing at random assumption. The robustness of this assumption was assessed in sensitivity analyses under the assumption of missingness not at random through use of pattern mixture models (appendix). To enhance efficiency and minimise bias in the imputation procedure, we included variables for which documentation was complete or almost complete even if they were irrelevant to substantive analysis. Covariates included in Model 1 were then analysed in models including all patients (Model 2) and restricted to non-severe cases (Model 3) through use of imputed datasets. For ease of clinical interpretation, particularly where risk factors were associated with high mortality, we converted odds ratios from logistic regression models to risk ratios. Accuracy of the three models was analysed by computing areas under the receiver operating characteristic (ROC) curve. For children with non-severe pneumonia, we calculated crude absolute risks of mortality by individual risk factor status as proportions with associated binomial exact 95% CIs. Subdivision of patients with non-severe pneumonia into groups by status of risk factor (with or without risk factor of interest) enabled us to compare risks of death between patients with and patients without the risk factor, and between those with non-severe pneumonia and an individual risk factor and patients with WHO-defined severe pneumonia. No allowance was made for multiple hypothesis testing. Results showing statistical significance close to p=0·05 should therefore be interpreted with caution. We analysed data with STATA version 12.0. The reporting of this study conforms to STROBE recommendations.

Role of the funding source
The funder of the study had no role in study design, data collection, data analysis, data interpretation, or writing of the manuscript. This research was funded by the Department of Health and Human Services, National Institutes of Health, National Institute of Allergy and Infectious Diseases (U01 AI083978). This manuscript was drafted with support from the Maine Department of Health and Human Services, Bureau of Preventive Health. Support for data collection, data management, data analysis, and data interpretation was provided by the International Centre for Diarrhoeal Disease Research, Bangladesh, and the Children’s Hospital of Philadelphia. The authors declare no competing interests. The funder of the study played no role in study design, data collection, data analysis, data interpretation, or writing of the manuscript.
the report. The corresponding author had full access to all data and had final responsibility for the decision to submit for publication.

Results
21832 records were available for children aged 2–59 months admitted to any one of 14 hospitals with a diagnosis of pneumonia between March 1, 2014, and Feb 29, 2016. Records for 5875 children with admission diagnoses that would exclude them from the WHO pneumonia case management algorithm were also excluded from the study. Children born before November, 2010, were excluded in keeping with our aim to study children in the post-vaccination era; therefore, analysis included 16162 children (figure 1).

The median age of the study population was 12 months (IQR 7–24) (table 2). Of 16127 children for whom sex was documented, 7180 (45%) were girls. 5313 (33%) of 16162 patients were admitted to hospitals in malaria endemic regions of Kenya. By use of documented clinical diagnosis and available information on clinical signs, we assigned pneumonia severity categories to patients on the basis of the WHO 2013 classification (table 1).

12025 (74%) of 16162 patients had non-severe pneumonia and 3468 (22%) of 16162 had severe pneumonia. 669 (4%) of 16162 children had inadequate documentation on key clinical signs required to assign their category of severity. Lower chest wall indrawing was present in 7514 (61%) of 12323 children, of whom 5320 (71%) were classified with non-severe pneumonia. We excluded children diagnosed with severe acute malnutrition at admission from analysis but calculated WAZ using WHO child growth standards for children remaining in the cohort because data for these two variables were complete for most patients studied. Although low WAZ might be due to stunting (chronic malnutrition), overlap exists across the categories of severity for mid-upper-arm circumference, WHZ, and WAZ. Each child was assigned to a nutritional category: normal WAZ (≥–2 SD), low WAZ (<–2 to –3), and very low WAZ (<–3). Data to calculate WAZ were available for 15300 children of whom 12324 (80%) had a normal score, 1861 (12%) had a low score, and 1145 (8%) had a very low score—none of the children were assigned a diagnosis of severe acute malnutrition by the admitting clinician. Pallor was present in 1257 (10%) of 12231 children with available data for the sign (table 2).

Documented outcomes were available for 16031 (99%) of 16162 children, and of those, 832 died (5.2%). 488 (14.2%) of 3434 children with severe pneumonia died compared with 322 (2.7%) of 11930 children within the non-severe category (risk ratio [RR] 5·3, 95% CI 4·6–6·0) (table 3). Mortality was higher in children younger than 12 months than those aged 12–59 months (RR 3·0, 2·6–3·4). Admission to hospitals in regions of high malaria prevalence was associated with increased risk of death (RR 1·3, 95% CI 1·2–1·5). Mortality also increased with decreasing WAZ. Among children with WAZ of <–2 SD or more, mortality was 3·8% whereas 7·6% of children with WAZ less than –2 SD to –3 SD and 11·2% of those with scores less than –3 SD died (score test for linear trend p<0·0001). Similarly, mortality increased proportionate to increasing severity of pallor and dehydration.

Other characteristics that were associated with increased mortality were elevated respiratory rate (>70 breaths per min), lower chest wall indrawing, axillary temperature (>39°C), and female sex. Immunisation status was only weakly associated with mortality (table 3).

Adjusted models included all variables analysed in univariate analyses except immunisation status. Parameter estimates from Model 1 (complete case analysis) were similar to those obtained in analyses in which missing data were imputed (table 4). In the full model in which imputed data were used (Model 2), severe pneumonia (>vs non-severe pneumonia) was independently associated with high mortality (adjusted risk ratio [aRR] 3·9, 95% CI 3·7–4·1). Other characteristics that were strongly associated with mortality (at least double

| Frequency or median | Number of patients |
|---------------------|--------------------|
| Age (months)       | 12 (7–22)          |
| Female             | 7180 (45%)         |
| Hospital location: high malaria prevalence* | 5313 (33%) |
| Immunisation status not up to date† | 373 (4%) |
| Pneumonia severity | 16162              |
| Non-severe         | 12 025 (74%)       |
| Severe             | 3468 (21%)         |
| Unclassified       | 669 (4%)           |
| Respiratory rate (breaths per min) | 10 992 |
| 2–11 months        | 56 (45–64)         |
| 12–59 months       | 48 (40–60)         |
| Lower chest wall indrawing present | 7514 (61%) |
| High fever (axillary temperature ≥39°C) | 2049 (18%) |
| WHZ                | 15 330             |
| >–2 SD (normal)    | 12 324 (80%)       |
| <–2 SD (low)       | 1861 (12%)         |
| <–3 SD (very low)  | 1145 (7%)          |
| Pallor             | 12 231             |
| Absent             | 10 974 (90%)       |
| Mild to moderate    | 874 (7%)           |
| Severe             | 383 (3%)           |
| Dehydration        | 16 091             |
| Absent or no dehydration | 14 038 (91%) |
| Some dehydration   | 902 (6%)           |
| Severe dehydration | 551 (3%)           |

Data are median (IQR) or n (%). WAZ=weight-for-age Z score. *Versus low to very low malaria prevalence. †Diphtheria, pertussis, tetanus, Hemophilus influenzae type b, hepatitis B, and pneumococcal vaccines: fewer than three doses at 3 months of age or older or fewer than two doses at 2 months of age or older.

Table 2: Demographic and clinical characteristics of the study participants
the risk of death) were mild to moderate (aRR 3.4–4.9, 95% CI 3.2–3.6) and severe (5.6, 5.1–6.1) pallor, lower chest wall indrawing (2.0–1.8–2.1), WAZ < –3 SD (2.1, 1.9–2.2), infants versus children aged 12–59 months (2.5–2.4–2.7), and children with severe dehydration (2.2, 2.0–2.4) versus no dehydration. Female sex, admission to hospital in a region of high malaria prevalence, elevated axillary temperature (≥39°C) or respiratory rate (≥70 breaths per min), and WAZ < –2 to < –3 SD were all also associated with increased mortality (aRR 1.3–1.9).

We further examined the independent associations for mortality within the category of non-severe pneumonia (for whom outpatient management is recommended) through use of imputed datasets (Model 3). The findings of this analysis were consistent with models using the full datasets (table 4). Areas under the ROC curve for each model showed good accuracy (0.8565, 0.8456, and 0.8115 for model 1, 2, and 3 respectively). Plots from the ROC analysis are provided in the appendix.

In subgroup analyses confined to children with non-severe pneumonia, estimated mortality of children with WAZ less than –3 SD (7.5%, 95% CI 4.9–11.8), mild to moderate pallor (7.8%, 5.1–11.4), and severe pallor (11.2%, 7.7–15.6) was similar to mortality observed in children with severe pneumonia (upper bound of 95% CI >10%). Risk of death for children with non-severe pneumonia and one risk factor—infants, admission to hospital in a malaria endemic region, respiratory rate of 70 breaths per min or higher, WAZ < –3 SD was mild to moderate (aRR 3.4, 95% CI 2.4–4.7) or some dehydration (2.2, 2.0–2.4) versus no dehydration. Female sex, admission to hospital in a region of high malaria prevalence, elevated axillary temperature (≥39°C) or respiratory rate (≥70 breaths per min), and WAZ < –2 to < –3 SD were all also associated with increased mortality (aRR 1.3–1.9).

We sought to determine the risk factors for pneumonia mortality in our study (5.2%), in a period after the introduction of conjugate vaccines, was higher than mortality reported in a previous systematic review25 that included infants younger than 2 months of age (pooled case fatality 3.9%). However, our results are similar to those from a secondary analysis of data from a clinical trial26 done in eight Kenyan hospitals before the introduction of the pneumococcal vaccine (overall pneumonia mortality 5.4%). The difference observed between the systematic review and clinical trial26 and our study might reflect higher baseline risk in sub-Saharan Africa, or selection of children with

### Table 3: Univariate associations for mortality among all children admitted to hospital with pneumonia

| Pneumonia classification | Number of deaths/number of patients | Mortality (%) | RR (95% CI) | p value* | P trend |
|--------------------------|-----------------------------------|--------------|-------------|---------|---------|
| Non-severe               | 322/11,930                        | 2.7%         | 1           | -       | -       |
| Severe                   | 488/14,344                        | 14.2%        | 5.3 (4.6–6.0) | <0.0001 | -       |
| Unclassified             | 22/667                            | 3.3%         | 1.2 (0.8–1.9) | <0.0001 | -       |
| Age                      |                                   |              |             |         |         |
| 12–59 months             | 228/8,471                         | 2.7%         | 1           | -       | -       |
| 2–11 months              | 604/7,560                         | 8.0%         | 3.0 (2.6–3.4) | <0.0001 | -       |
| Sex                      |                                   |              |             |         |         |
| Male                     | 380/8,767                         | 4.3%         | 1           | -       | -       |
| Female                   | 447/7,715                         | 6.3%         | 1.4 (1.3–1.7) | <0.0001 | -       |
| Malaria prevalence       |                                   |              |             |         |         |
| Low                      | 504/10,747                        | 4.7%         | 1           | -       | -       |
| High                     | 328/5,284                         | 6.2%         | 1.3 (1.2–1.5) | <0.0001 | -       |
| Immunisation status      |                                   |              |             |         |         |
| Up to date               | 404/8,683                         | 4.7%         | 1           | -       | -       |
| Not up to date           | 27/365                            | 7.4%         | 1.6 (1.1–2.3) | 0.04 | -       |
| Respiratory rate         |                                   |              |             |         |         |
| <70 breaths per min      | 125/12,54                         | 4.1%         | 1           | -       | -       |
| ≥70 breaths per min      | 394/9,657                         | 10.0%        | 2.4 (2.0–3.0) | <0.0001 | -       |
| Lower chest wall indrawing|                                   |              |             |         |         |
| Absent                   | 132/4,770                         | 2.8%         | 1           | -       | -       |
| Present                  | 468/7,455                         | 6.3%         | 2.3 (1.9–2.7) | <0.0001 | -       |
| Axillary temperature     |                                   |              |             |         |         |
| <39°C                    | 394/9,587                         | 4.3%         | 1           | -       | -       |
| ≥39°C                    | 158/2,026                         | 7.8%         | 1.9 (1.6–2.3) | <0.0001 | -       |
| WAZ                      |                                   |              |             |         |         |
| ≥2 SD                    | 469/12,235                        | 3.8%         | 1           | -       | <0.0001|
| <2 up to –3 SD           | 140/18,43                         | 7.6%         | 2.0 (1.6–2.3) | <0.0001 | -       |
| <3 SD                    | 127/11,334                        | 11.2%        | 2.9 (2.4–3.4) | <0.0001 | -       |
| Pallor                   |                                   |              |             |         |         |
| Absent                   | 396/10,888                        | 3.6%         | 1           | -       | <0.0001|
| Mild to moderate         | 127/8,655                         | 14.7%        | 4.0 (3.3–4.7) | <0.0001 | -       |
| Severe                   | 68/3,182                          | 17.8%        | 4.8 (3.8–6.0) | <0.0001 | -       |
| Dehydration              |                                   |              |             |         |         |
| Absent                   | 615/14,526                        | 4.4%         | 1           | -       | <0.0001|
| Some                     | 59/8,902                          | 6.6%         | 1.5 (1.2–1.9) | <0.0001 | -       |
| Severe                   | 126/5,443                         | 23.2%        | 5.2 (4.4–6.0) | <0.0001 | -       |

Weight-for-age Z score (WAZ) based on WHO reference growth standards. Immunisation status omitted in multivariable models. RR=risk ratio. *p values derived from χ² test.

**Discussion**

We sought to determine the risk factors for pneumonia mortality among children admitted to hospital in settings with high prevalence of comorbidities. By contrast with previous studies in highly controlled research environments,20 we proposed to study children who were managed in real-life settings in which most studies on guideline adherence suggest only partial compliance.21–24 This study represents the largest published individual analysis of risk factors for mortality among children with WHO-defined non-severe pneumonia. Pneumonia mortality in our

www.thelancet.com/lancetgh Vol 6 January 2018
Articles

| Pneumonia classification | Model 1: complete case analysis (all pneumonia cases) | Model 2: multiple imputation analysis (all pneumonia cases) | Model 3: multiple imputation analysis (non-severe pneumonia only) |
|--------------------------|-----------------------------------------------------|---------------------------------------------------------|-------------------------------------------------------------|
|                          | Adjusted 95% CI                                     | Adjusted 95% CI                                        | Adjusted 95% CI                                              |
| Non-severe               | 1 (ref) -                                          | 1 (ref) -                                               | -                                                          |
| Severe                   | 4.2 (3.4–5.2)                                      | 3.9 (3.7–4.1)                                          | -                                                          |

| Age                      | Model 1: complete case analysis (all pneumonia cases) | Model 2: multiple imputation analysis (all pneumonia cases) | Model 3: multiple imputation analysis (non-severe pneumonia only) |
|--------------------------|-----------------------------------------------------|---------------------------------------------------------|-------------------------------------------------------------|
| 12-59 months             | 1 (ref) -                                          | 1 (ref) -                                               | -                                                          |
| 2-11 months              | 2.7 (2.2–3.4)                                      | 2.5 (2.4–2.7)                                          | 2.8 (2.6–3.1)                                              |

| Sex                      | Model 1: complete case analysis (all pneumonia cases) | Model 2: multiple imputation analysis (all pneumonia cases) | Model 3: multiple imputation analysis (non-severe pneumonia only) |
|--------------------------|-----------------------------------------------------|---------------------------------------------------------|-------------------------------------------------------------|
| Male                     | 1 (ref) -                                          | 1 (ref) -                                               | -                                                          |
| Female                   | 1.6 (1.3–1.9)                                      | 1.5 (1.4–1.6)                                          | 1.4 (1.3–1.6)                                              |

| Malaria prevalence       | Model 1: complete case analysis (all pneumonia cases) | Model 2: multiple imputation analysis (all pneumonia cases) | Model 3: multiple imputation analysis (non-severe pneumonia only) |
|--------------------------|-----------------------------------------------------|---------------------------------------------------------|-------------------------------------------------------------|
| Low                      | 1 (ref) -                                          | 1 (ref) -                                               | -                                                          |
| High                     | 1.3 (1.1–1.7)                                      | 1.3 (1.2–1.4)                                          | 1.6 (1.4–1.7)                                              |

| Respiratory rate         | Model 1: complete case analysis (all pneumonia cases) | Model 2: multiple imputation analysis (all pneumonia cases) | Model 3: multiple imputation analysis (non-severe pneumonia only) |
|--------------------------|-----------------------------------------------------|---------------------------------------------------------|-------------------------------------------------------------|
| <70 breaths per min      | 1 (ref) -                                          | 1 (ref) -                                               | -                                                          |
| ≥70 breaths per min      | 1.8 (1.4–2.3)                                      | 1.8 (1.7–1.9)                                          | 2.3 (2.1–2.6)                                              |

| Lower chest wall indrawing | Model 1: complete case analysis (all pneumonia cases) | Model 2: multiple imputation analysis (all pneumonia cases) | Model 3: multiple imputation analysis (non-severe pneumonia only) |
|---------------------------|-----------------------------------------------------|---------------------------------------------------------|-------------------------------------------------------------|
| Absent                    | 1 (ref) -                                          | 1 (ref) -                                               | -                                                          |
| Present                   | 1.8 (1.4–2.4)                                      | 2.0 (1.8–2.1)                                          | 2.5 (2.2–2.7)                                              |

| Axillary temperature      | Model 1: complete case analysis (all pneumonia cases) | Model 2: multiple imputation analysis (all pneumonia cases) | Model 3: multiple imputation analysis (non-severe pneumonia only) |
|---------------------------|-----------------------------------------------------|---------------------------------------------------------|-------------------------------------------------------------|
| <39°C                     | 1 (ref) -                                          | 1 (ref) -                                               | -                                                          |
| ≥39°C                     | 1.9 (1.5–2.3)                                      | 1.8 (1.7–1.9)                                          | 1.9 (1.7–2.1)                                              |

| WAZ                       | Model 1: complete case analysis (all pneumonia cases) | Model 2: multiple imputation analysis (all pneumonia cases) | Model 3: multiple imputation analysis (non-severe pneumonia only) |
|---------------------------|-----------------------------------------------------|---------------------------------------------------------|-------------------------------------------------------------|
| ≥-2 SD                    | 1 (ref) -                                          | 1 (ref) -                                               | -                                                          |
| <-2 to -3 SD              | 1.8 (1.4–2.4)                                      | 1.90 (1.8–2.0)                                         | 2.4 (2.2–2.7)                                              |
| <-3 SD                    | 2.1 (1.6–2.9)                                      | 2.05 (1.9–2.2)                                         | 3.8 (3.4–4.3)                                              |

| Pallor                    | Model 1: complete case analysis (all pneumonia cases) | Model 2: multiple imputation analysis (all pneumonia cases) | Model 3: multiple imputation analysis (non-severe pneumonia only) |
|---------------------------|-----------------------------------------------------|---------------------------------------------------------|-------------------------------------------------------------|
| Absent                    | 1 (ref) -                                          | 1 (ref) -                                               | -                                                          |
| Mild to moderate           | 3.5 (2.7–4.5)                                      | 3.38 (3.2–3.6)                                         | 3.4 (3.0–3.8)                                              |
| Severe                    | 6.1 (4.3–8.7)                                      | 5.58 (5.1–6.1)                                         | 5.9 (5.1–6.8)                                              |

| Dehydration               | Model 1: complete case analysis (all pneumonia cases) | Model 2: multiple imputation analysis (all pneumonia cases) | Model 3: multiple imputation analysis (non-severe pneumonia only) |
|---------------------------|-----------------------------------------------------|---------------------------------------------------------|-------------------------------------------------------------|
| Absent                    | 1 (ref) -                                          | 1 (ref) -                                               | -                                                          |
| Some                      | 2.5 (1.7–3.6)                                      | 2.17 (2.0–2.4)                                         | 2.2 (2.0–2.6)                                              |
| Severe                    | 2.3 (1.6–3.1)                                      | 2.21 (2.0–2.4)                                         | No observations                                             |

Weight-for-age Z score (WAZ) classification based on WHO reference growth standards. RR-risk ratio. *Models adjusted for age, sex (a priori covariates), hospital location (fixed effect), immunisation status, WHO pneumonia category (except where only patients with non-severe pneumonia are analysed), respiratory rate, temperature, WAZ, pallor, and dehydration.

Table 4: Multivariable models for risk factors for mortality—complete case analysis and multiple imputation analyses (all pneumonia cases and non-severe pneumonia only)

Low risk in prospective studies included in the systematic review. In our study, less than 0·5% of children with non-severe pneumonia and without any of the risk factors studied died. Mortality among non-severe cases (as defined by WHO) in our study was also higher (2.7%) than other studies that report mortality rates less than 1%.

WHO guidelines recognise severe malnutrition defined by WHZ or mid-upper-arm circumference as a risk factor for mortality, warranting inpatient care when present in association with pneumonia. The use of WAZ is no longer recommended as a screening approach to identify severe acute malnutrition. We observed a high prevalence of low to very low WAZ (almost 20% of the study population), even after exclusion of cases of severe acute malnutrition were identified by the admitting clinician on the basis of mid upper-arm-circumference, WHZ, visible severe wasting, or oedema due to kwashiorkor. Children with non-severe pneumonia and WAZ less than –3 SD and without severe acute malnutrition had a risk of death almost three times higher than patients with non-severe pneumonia. Increased risk of death among children with poor nutritional status is widely thought to be linked to deficiencies in immune function. Consistent with our findings, a systematic review of children with pneumonia and malnutrition (defined by use of WAZ or WHZ) reported invariably high risks of death for both moderate or severe forms of malnutrition in all 16 studies included.

The presence of pallor, even when classified as mild or moderate, was also strongly associated with mortality across all our analyses. Severe pallor is commonly used as a clinical marker for anaemia in settings where laboratory diagnostics are unavailable, with sensitivity and specificity above 80% for children with packed cell volumes less than 15%. Anaemia is a common presenting feature in sub-Saharan African children, manifesting acutely in conditions such as malaria and chronically because of inadequate nutritional iron intake or helminthic infestation. Severe anaemia has been associated with increased mortality in previous studies of children with pneumonia, our study presents new evidence showing risk for mild forms.

These findings have important implications for the WHO policy advocating for outpatient treatment for all but the children with a small set of clinical signs that define severe pneumonia. Although severe pallor would generally warrant admission to hospital, the existing recommendations for case management do not consider very low WAZ or mild to moderate pallor as important risk factors. Thus 1272 (11%) of 12 025 children defined as having non-severe pneumonia in this study with either of these two risk factors would be expected to be managed at home.

A large subpopulation of children had at least one risk factor that was independently associated with mortality, and absolute risks of death higher than the upper confidence limit of that for all non-severe pneumonia cases, but below that for severe pneumonia (mortality between 3% and 10%). The characteristics falling in this category were age younger than 12 months, admission to hospital in a location with high malaria prevalence, elevated respiratory rate (≥70 breaths per min), WAZ less than –2 to –3 SD, lower chest wall indrawing, and moderate dehydration. Our analysis concurs with the findings of a systematic review of risk factors for mortality in children with pneumonia (almost entirely
done in the pre-pneumococcal vaccine period). In this review that included over 135,000 children from sub-Saharan Africa, the risk factors for mortality in common with our study were low WAZ, female sex, and age younger than 12 months. Infants, who constituted half the population in our study and had mortality three times greater than children aged 12–59 months, are also not assigned priority under the WHO guidelines despite urgent recommendations for review of the guidelines after previous studies. WHO modified the pneumonia case management algorithm to define only two levels of severity in place of the previous three. This recommendation might simplify training of health workers at the primary care and community level by reducing decision making to whether a child requires inpatient or outpatient care. The downgrading of lower chest wall indrawing, previously a sign defining the need for admission, has been challenged in settings of high mortality, where, among other signs, it is associated with fatal pneumonia. Our analyses show the presence at hospital level of many children with signs of unrecognised intermediate severity, including lower chest wall indrawing. At least in Kenya, such children appear to have an in-hospital mortality of at least 3%.

Although we highlight associations with mortality in this article, identification of characteristics that might optimise decision making on admission (linked to risk of poor outcome) while being parsimonious to promote simplicity will require additional analyses. In a related article, we attempt to use machine learning modelling techniques to rank risk factors for pneumonia mortality and use decision curve analysis to explore potential net benefit of the use of highest-ranking risk factors in the decision to admit non-severe pneumonia cases.

Although patients with severe pneumonia were largely prescribed guideline-recommended broad-spectrum antibiotics on admission (unpublished), mortality was more than five times higher than that observed for the non-severe category. The high mortality in this group might suggest late clinical presentation (justifying the need for interventions targeting improved care-seeking or referral structures), or inadequacy of existing treatments, suggesting the need for research to explore alternative antibiotic regimens or interventions to improve supportive care in settings with scarce resources. Supportive care might include improved use of routine pulse oximetry, use of reliable oxygen supplies, appropriate administration of fluids and feeding, and optimisation of nursing care.

---

**Figure 2: Estimated risks of death among patients with non-severe pneumonia by risk factor status**

WAZ—weight-for-age Z score. Excludes patients aged 2–11 months, girls, treatment in hospital in area with high prevalence of malaria, immunisations not up to date, respiratory rate ≥70 breaths per min, chest indrawing present, axillary temperature ≥39°C, WAZ <–2 SD, mild to moderate or severe pallor, or moderate dehydration.

| Risk Factor | Mortality (95% CI) |
|-------------|-------------------|
| All pneumonia | 5.2% (4.9–5.5) |
| All non-severe | 2.7% (2.4–3.0) |
| Non-severe, no risk factors* | 0.2% (0.0–0.9) |
| Age 12–59 months | 1.5% (1.2–1.8) |
| Age 2–11 months | 4.2% (3.7–4.8) |
| Male sex | 2.2% (1.9–2.6) |
| Female sex | 3.3% (2.8–3.8) |
| Low malaria prevalence | 2.2% (1.9–2.5) |
| High malaria prevalence | 3.8% (3.2–4.4) |
| Immunisation up to date | 2.3% (1.9–2.7) |
| Immunisation not up to date | 4.2% (3.0–7.6) |
| Respiratory rate ≥70 breaths per min | 1.8% (1.6–2.2) |
| Respiratory rate ≤70 breaths per min | 4.1% (3.5–4.7) |
| Chest indrawing absent | 1.5% (1.1–1.9) |
| Chest indrawing present | 3.2% (2.7–3.7) |
| Axillary temperature <39°C | 2.0% (1.7–2.3) |
| Axillary temperature ≥39°C | 3.8% (2.8–5.4) |
| WAZ ≥–2 SD | 1.8% (1.5–2.1) |
| WAZ <–2 to –3 SD | 4.4% (3.4–5.6) |
| WAZ <–3 SD | 7.5% (4.9–11.8) |
| Pallor absent | 1.8% (1.5–2.1) |
| Mild to moderate pallor | 7.8% (5.1–11.4) |
| Severe pallor | 11.2% (7.7–15.6) |
| Dehydration absent | 2.5% (2.2–2.8) |
| Moderate dehydration | 5.1% (3.6–6.9) |
The large sample size taken from multiple hospitals across the country resulted in precise estimates that are broadly representative of the population of children admitted to hospital with pneumonia in many district hospitals in sub-Saharan Africa. The collection of data covering 2 years further enhanced representativeness through elimination of seasonal bias—an important consideration in studies on acute respiratory infections in children. Our analysis is based on clinical features recorded by many junior clinicians providing routine care; therefore, the interpretation of clinical signs is likely to vary. However, non-differential misclassification resulting from variations in interpretation of signs would be expected to increase random error and reduce the precision of risk estimates. Arguably, inclusion of data captured by many clinicians in routine practice increases the generalisability of our findings.

Analysis is based on documentation of clinical assessments and subsequent data entry. This method has the potential for selection bias arising from misplaced patient records not included in the analysis. However, considerable effort was made to ensure comprehensive sampling through data entry immediately after the end of the inpatient stay with regular comparisons of clinical records retrieved against hospital admission registers. Rigorous training and close supervision of data clerks, use of structured patient admission forms with feedback on their completeness, and dissemination of standard clinical guidelines to health workers at the study hospitals improved the quality of data collected. We also applied multiple imputation to maximise the use of available data and did sensitivity analyses that supported the assumption that missingness was not associated with severity.

Inadequate individual data on malaria status led us to use hospital location (high vs low to very low malaria endemicity) as a crude proxy for this diagnosis. Overlap in clinical presentation of malaria from pneumonia might have undermined the validity of risk factor analyses outside high malaria endemic settings. To explore this possibility, we did subanalyses restricted to patients from very low malaria endemic sites (unpublished), which yielded parameter estimates that were consistent with those obtained through use of the full dataset including statistically significant (p<0.0001) associations with mortality for high fever and pallor (two signs common in malaria).

Systematic reviews have shown substantially lower risks of mortality among immunised children than unimmunised children: this association was not apparent in our study. A possible explanation for this absence might be errors of misclassification perhaps attributable to unreliable caregiver-reported data on immunisation status. Inclusion of data on oxygen saturation would have been potentially useful for further definition of risk; however, pulse oximetry is still not commonly available in the study hospitals. Confirmation of the extent to which treatments prescribed such as antibiotics, oxygen, and fluids were received was not possible.

We were unable to collect information on clinical characteristics and outcomes of children with pneumonia managed as outpatients. These data are not available from routine settings in hospitals that contributed data to this analysis, nor any other setting in sub-Saharan Africa. Unobserved factors might have prompted admission of some children—ie, clinicians might have been responding to so-called gut feeling about severity. Such clinical decisions might have resulted in a higher mortality among children admitted to hospital with a given profile of specified risk factors than among children not admitted to hospital with those same risk factors. However, even if risk of death among patients admitted to hospital with risk factors was higher than the risk among patients not admitted to hospital with the same risk factors, the potential for adverse consequences of implementation of the WHO guidelines in a country such as Kenya cannot be ignored. Thus, discharging of children at high risk would probably result in worse outcomes if hospital admission is effective. The only reason for promulgation of a guideline is to change practice, and in this case, that would mean admission of a reduced proportion of patients shown by our study to have a high risk of death.

Most children admitted to hospital in a period immediately preceding the implementation of policy guidelines that would classify them as having non-severe pneumonia presented with at least one factor associated with a risk of inpatient death above 3%. Among these children, mortality was substantially higher for those presenting with WAZ less than –3 SD or pallor (both mild to moderate and severe) than those without these risk factors. Additionally, infants, admission to hospital in a location with high malaria prevalence, WAZ less than −2 to −3 SD, females, elevated respiratory rate (≥70 breaths per min) or axillary temperature of 39°C or above, lower chest wall indrawing, and some dehydration were independently associated with increased mortality. Our study therefore identifies subgroups of patients at risk for future intervention studies to refine pneumonia case management among inpatients and outpatients. Our findings suggest that presence of WAZ less than −3 SD or any degree of pallor among children with WHO-defined non-severe pneumonia should be considered alongside the WHO criteria for admission care in contemporary African populations—a finding warranting further study. Although this outcome does not constitute formal evidence that making use of these factors would save lives, these results imply a risk of offering outpatient treatment to specific patient subgroups with non-trivial risks of mortality, and provide a foundation for future work to derive a simple risk score for implementation in clinical settings where WHO case management guidelines are in use.

Contributors
AA, RJL, and ME conceived the study and AA did the analyses with support from RJL and ME. AA drafted the initial manuscript with
We thank the Ministry of Health which gave permission for this work to be developed and has supported the implementation of the Clinical Information Network, together with the county health executives and all hospital management teams. Collaboration with officers from the Ministry of Health’s National Health Management Information System, the Monitoring and Evaluation Unit, and the Neonatal, Child and Adolescent Health Unit has been important to the initiation of the Clinical Information Network. We are grateful to the Kenya Paediatric Association for promoting the aims of the Clinical Information Network and the support they provide through their officers and membership. We also thank the paediatricians and clinical teams on all the paediatric wards who provide care to the children for whom this project is designed. Statistical support for the analyses involving multiple imputation was provided by David Gathara and Lucas Malia. This work was published with the permission of the Director of KEMRI. This study was supported by funds from a senior research fellowship awarded to ME by the Wellcome Trust. AA was supported by funds from a Wellcome Trust Strategic Award, the Initiative to Develop African Research Leaders (iDeA) Wellcome Trust award, and a Wellcome Trust core grant awarded to the KEMRI-Wellcome Trust Research Programme. RJL is supported by funds from the National Institute for Health Research Collaboration Awards. Clinical Information Network authors contributed to the design of the data collection tools, conduct of the work, collection of data, and data quality assurance that form the basis of this Article, and saw and approved the Article’s findings. All coauthors reviewed and approved the final version of the manuscript.

Declaration of interests

We declare no competing interests.

Acknowledgements

The Ministry of Health’s National Health Management Information System, Geneva: World Health Organization, 2013.

We declare no competing interests.

References

1 WHO. Pocket book of hospital care for children: guidelines for the management of common illnesses with limited resources, 2nd edn. Geneva: World Health Organization, 2013.
2 Scott JA, English M. What are the implications for childhood pneumonia of successfully introducing Hib and pneumococcal vaccines in developing countries? PLoS Med 2008; 5: e86.
3 WHO, UNICEF, UNAIDS, UN Population Fund. Towards the elimination of mother-to-child transmission of HIV: report of a WHO technical consultation, 9–11 November 2010, Geneva, Switzerland. Geneva: World Health Organization, 2011.
4 WHO. Guidelines: update on the management of severe acute malnutrition in infants and children. Geneva: World Health Organization, 2013.
5 Mulholland EK, Simoes EA, Costales MO, McGrath EJ, Manalac EM, Gove S. Standardized diagnosis of pneumonia in developing countries. Pediatr Infect Dis J 1992; 11: 77–81.
6 Agwem A, Opiyo N, English M. Experience developing national evidence-based clinical guidelines for childhood pneumonia in a low-income setting—making the GRADE? BMC Pediatr 2012; 12: 1.
7 Mulholland K, Carlin JB, Duke T, Weber M. The challenges of trials of antibiotics for pneumonia in low-income countries. Lancet Respir Med 2014; 2: 952–54.
8 Ayepeko P, Ogero M, Makone B, et al. Characteristics of admissions and variations in the use of basic investigations, treatments and outcomes in Kenyan hospitals within a new Clinical Information Network. Arch Dis Child 2016; 101: 223–29.
9 Kenya National Bureau of Statistics (KNBS) and ICF Macro. Kenya Demographic Health Survey 2014. Calverton: KNBS and ICF Macro, 2014.
10 Tutu T, Bitok M, Paton C, et al. Innovating to enhance clinical data management using non-commercial and open source solutions across a multi-center network supporting inpatient pediatric care and research in Kenya. J Am Med Inform Assoc 2016; 23: 184–92.
11 Health Services Unit KEMRI-Wellcome Trust Research Programme. Improving the delivery of hospital care in Africa. http://idoc-africa.org/ (accessed Jan 16, 2017).
12 Harris PA, Taylor R, Thielke R, Payne J, Gonzalez N, Conde JG. Research electronic data capture (REDCap)—a metadata-driven methodology and workflow platform for providing translational research informatics support. J Biomed Inform 2009; 42: 377–81.
13 Azur MJ, Stuart EA, Franjakis C, Leaf PJ. Multiple imputation by chained equations: what is it and how does it work? J Methods Psychiatr Res 2011; 20: 40–49.
14 Bell ML, Fairclough DL. Practical and statistical issues in missing data for longitudinal patient-reported outcomes. Stat Methods Med Res 2014; 23: 448–70.
15 Collins LM, Schafer JL, Kam CM. A comparison of inclusive and restrictive strategies in modern missing data procedures. Psychol Methods 2001; 6: 330–51.
16 Zhang J, Yu KF. What’s the relative risk? A method of correcting the odds ratio in cohort studies of common outcomes. JAMA 1998; 280: 1690–91.
17 von Elm E, Altman DG, Egger M, et al. The Strengthening the Reporting of Observational Studies in Epidemiology (STROBE) statement: guidelines for reporting observational studies. Lancet 2007; 370: 1453–72.
18 WHO, WHO Child Growth Standards. Geneva: World Health Organization, 2006.
19 Berkley J, Mwangi I, Griffiths K, et al. Assessment of severe malnutrition among hospitalized children in rural Kenya: comparison of weight for height and mid upper arm circumference. JAMA 2005; 294: 591–97.
20 Fox MP, Thea DM, Sadr-Hosseini S, et al. Low rates of treatment failure in children aged 2–59 months treated for severe pneumonia: a multisite pooled analysis. Clin Infect Dis 2013; 56: 978–87.
21 English M, Esamai F, Wasunna A, et al. Assessment of inpatient paediatric care in first referral level hospitals in 13 districts in Kenya. Lancet 2004; 363: 1948–53.
22 Hoque DM, Rahman M, Billah SM, et al. An assessment of the quality of care for children in eighteen randomly selected district and sub-district hospitals in Bangladesh. BMC Pediatr 2012; 12: 197.
23 Nolan T, Angos P, Cunha AJ, et al. Quality of hospital care for seriously ill children in less-developed countries. Lancet 2001; 357: 306–10.
24 Reyburn H, Mwakasungula E, Chonya S, et al. Clinical assessment and treatment in paediatric wards in the north-east of the United Republic of Tanzania. Bull World Health Organ 2008; 86: 132–39.
25 Nair H, Simoe EA, Rudan I, et al. Global and regional burden of hospital admissions for severe acute lower respiratory infections in young children in 2010: a systematic analysis. Lancet 2013; 381: 1380–90.
26 Ayepeko P, Okiro EA, Edwards T, Nyamai R, English M. Variations in mortality in children admitted with pneumonia to Kenyan hospitals. PLoS One 2012; 7: e46722.
27 Ryttter MJ, Kolte L, Briand A, Fris H, Christensen VB. The immune system in children with malnutrition—a systematic review. PLoS One 2014; 9: e105017.
28 Waterlow JC, Alleyne GA. Protein malnutrition in children: advances in knowledge in the last ten years. Adv Protein Chem 1979; 25: 117–241.
29 Tomkins AM, Garlick PJ, Schofield WN, Waterlow JC. The combined effects of infection and malnutrition on protein metabolism in children. Clin Sci (Lond) 1983; 65: 313–24.
30 Chistia MJ, Tehruigge M, La Vincenzo S, Graham SM, Duke T. Pneumonia in severely malnourished children in developing countries—mortality risk, aetiology and validity of WHO clinical signs: a systematic review. Trop Med Int Health 2009; 14: 1273–89.
31 Zucker JR, Perkins BA, Jafari H, Otieno J, Oborony C, Campbell CC. Clinical signs for the recognition of children with moderate or severe anaemia in western Kenya. Bull World Health Organ 1997; 75 (suppl 1): 97–102.
32 Weber MW, Kellingray SD, Palmer A, Jaffar S, Mulholland EK, Greenwood BM. Pallor as a clinical sign of severe anaemia in children: an investigation in the Gambia. Bull World Health Org 1997; 75(suppl 1): 113–18.

33 Jroundi I, Mahraoui C, Benmessaoud R, et al. Risk factors for a poor outcome among children admitted with clinically severe pneumonia to a university hospital in Rabat, Morocco. Int J Infect Dis 2014; 28: 164–70.

34 Enarson PM, Gie RP, Mwansambo CC, et al. Potentially modifiable factors associated with death of infants and children with severe pneumonia routinely managed in district hospitals in Malawi. PLoS One 2015; 10: e0133365.

35 Moschovis PP, Banajeh S, MacLeod WB, et al. Childhood anemia at high altitude: risk factors for poor outcomes in severe pneumonia. Pediatrics 2013; 132: e1156–62.

36 Sonigo M, Pellegrin MC, Becker G, Luzzerini M. Risk factors for mortality from acute lower respiratory infections (ALRI) in children under five years of age in low and middle-income countries: a systematic review and meta-analysis of observational studies. PLoS One 2015; 10: e0136380.

37 Jeena P, Thea DM, MacLeod WB, et al. Failure of standard antimicrobial therapy in children aged 3–59 months with mild or asymptomatic HIV infection and severe pneumonia. Bull World Health Org 2006; 84: 269–75.

38 McNally J, Jeena PM, Gajee K, et al. Effect of age, polymicrobial disease, and maternal HIV status on treatment response and cause of severe pneumonia in South African children: a prospective descriptive study. Lancet 2007; 369: 1446–51.

39 Reed C, Madhi SA, Khugman KP, et al. Development of the Respiratory Index of Severity in Children (RISC) score among young children with respiratory infections in South Africa. PLoS One 2012; 7: e27793.

40 Tuti T, Agewuyu A, Mwaniki P, Peek N, English M. An exploration of mortality risk factors in non-severe pneumonia in children using clinical data from Kenya. BMC Med 2017; 15: 201.

41 Gathara D, Nyamai R, Were F, et al. Moving towards routine evaluation of quality of inpatient pediatric care in Kenya. PLoS One 2015; 10: e0137048.

42 Enoch AJ, English M, Shepperd S. Does pulse oximetry use impact health outcomes? A systematic review. Arch Dis Child 2016; 101: 694–700.

43 Moschovis PP, Hibberd PL. Pulse oximetry: an important first step in improving health outcomes, but is of little use if there is no oxygen. Arch Dis Child 2016; 101: 685.

44 Ampofo K, Bender J, Sheng X, et al. Seasonal invasive pneumococcal disease in children: role of preceding respiratory viral infection. Pediatrics 2008; 122: 229–37.

45 Tuti T, Bitok M, Malla I, et al. Improving documentation of clinical care within a clinical information network: an essential initial step in efforts to understand and improve care in Kenyan hospitals. BMJ Glob Health 2016; 1: e000028.

46 Bassat Q, Machevo S, O’Callaghan-Gordo C, et al. Distinguishing malaria from severe pneumonia among hospitalized children who fulfilled integrated management of childhood illness criteria for both diseases: a hospital-based study in Mozambique. Am J Trop Med Hyg 2011; 85: 626–34.

47 English M, Punt J, Mwangi I, McHugh K, Marsh K. Clinical overlap between malaria and severe pneumonia in African children in hospital. Trans R Soc Trop Med Hyg 1996; 90: 658–62.

48 Kallander K, Nsungwa-Sabiiti J, Peterson S. Symptom overlap for malaria and pneumonia—policy implications for home management strategies. Acta Trop 2004; 90: 211–14.

49 Van den Bruel A, Thompson M, Buntinx F, Mant D. Clinicians’ gut feeling about serious infections in children: observational study. BMJ 2012; 345: e6144.