High burden and seasonal variation of paediatric scabies and pyoderma prevalence in The Gambia: A cross-sectional study

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Abstract

Background
Scabies is a WHO neglected tropical disease common in children in low- and middle-income countries. Excoriation of scabies lesions can lead to secondary pyoderma infection, most commonly by Staphylococcus aureus and Streptococcus pyogenes (group A streptococcus, GAS), with the latter linked to acute post-streptococcal glomerulonephritis (APSGN) and potentially rheumatic heart disease (RHD). There is a paucity of data on the prevalence of these skin infections and their bacterial aetiology from Africa.

Methodology/Principal findings
A cross-sectional study, conducted over a four-month period that included the dry and rainy season, was conducted to determine the prevalence of common skin infections in Sukuta, a peri-urban settlement in western Gambia, in children <5 years. Swabs from pyoderma lesions were cultured for S. aureus and GAS. Of 1441 children examined, 15.9% had scabies (95% CI 12.2–20.4), 17.4% had pyoderma (95% CI 10.4–27.7) and 9.7% had fungal infections (95% CI 6.6–14.0). Scabies was significantly associated with pyoderma (aOR 2.74, 95% CI 1.61–4.67). Of 250 pyoderma swabs, 80.8% were culture-positive for S. aureus, and 50.8% for GAS. Participants examined after the first rains were significantly more likely to have pyoderma than those examined before (aRR 2.42, 95% CI 1.38–4.23), whereas no difference in scabies prevalence was seen (aRR 1.08, 95% CI 0.70–1.67). Swab positivity was not affected by the season.

Conclusions/Significance
High prevalence of scabies and pyoderma were observed. Pyoderma increased significantly during the rainy season. Given the high prevalence of GAS pyoderma among children, further research on the association with RHD in West Africa is warranted.
Scabies is a WHO neglected tropical disease which is common in low- and middle-income countries with tropical climates, which can lead to serious complications via secondary bacterial infection of skin lesions. Group A streptococcal skin infection is known to cause kidney disease, but may also contribute to rheumatic heart disease. Despite this, little is known about the prevalence of common paediatric skin infections in sub-Saharan Africa, or which bacteria most commonly cause skin infections. We conducted a cross-sectional study of skin infections in The Gambia into the prevalence of scabies, pyoderma and fungal infections in children <5 years. We examined 1441 children and found a scabies, pyoderma and fungal infection prevalence of 15.9%, 17.4% and 9.7% respectively, with significantly increased presence of bacterial skin infections in the rainy season. Over half of bacterial skin infections were found to be positive for group A streptococcus. Our study shows that scabies and pyoderma occur commonly in young children in The Gambia, representing a significant exposure to group A streptococcus at a young age, and that there is a strong seasonal effect on pyoderma, which may have implications for treatment strategies.

Introduction

Streptococcus pyogenes (group A streptococcus, GAS) is a pathogen that causes a wide spectrum of disease, from superficial skin infections through to invasive sepsis and streptococcal toxic shock syndrome [1]. It continues to cause a significant burden of morbidity and mortality globally, particularly in low- and middle-income countries (LMIC) [2]. It is also associated with autoimmune-mediated, post-infective sequelae including acute post-streptococcal glomerulonephritis (APSGN) and acute rheumatic fever (ARF) leading to chronic rheumatic heart disease (RHD) [3]. RHD has largely disappeared from high-income countries except in neglected groups such as Aboriginal Australians [3, 4], but remains a significant problem in LMIC including those in sub-Saharan Africa (sSA) [5]. The classical understanding of ARF involves an acute GAS pharyngitis infection with specific “rheumatogenic” GAS emm serotypes [6, 7]. It is, however, increasingly thought that this explanation is incomplete, with a diversity of emm types and GAS skin infections potentially playing an important role in LMIC [8, 9]. This is supported by data from countries where GAS pyoderma and RHD are both highly prevalent, but pharyngitis and GAS pharyngeal carriage are low [3, 10, 11]. A significant proportion of GAS skin infections are attributable to secondary bacterial infection of scabies lesions, a WHO neglected tropical disease, known to be particularly prevalent in young children living in poverty-related conditions in tropical countries [12–14]. Despite this, there are few data sources on scabies epidemiology for sSA [15–20].

Pyoderma (defined as any infection of the skin involving pus) is common in LMIC independently of its association with scabies [12], with over 162 million children estimated to be suffering from impetigo globally [21]. Epidemiological and microbiological data on pyoderma in Africa is scarce [3, 22–24]. Due to the importance of GAS as a pathogen in sSA and M-protein type-specific vaccines in development, a registry of GAS infections in Africa has been initiated to address the paucity of microbiological data [25]. Superficial fungal infections are also common globally, particularly in children in LMIC. The burden in Africa is heterogeneous, with prevalence in schoolchildren ranging from 10–80% [26].
In The Gambia, the prevalence of these common skin infections and the bacterial aetiology of pyoderma in children are unknown. To provide preliminary data, we performed a cross-sectional study to determine prevalence of scabies, pyoderma and fungal skin infections in children aged <5 years in a peri-urban community in western Gambia. We also aimed to characterise the microbiological aetiology of pyoderma in this setting.

**Methods**

**Setting**

The Gambia is the smallest country by area in mainland Africa, with a population of 1.9 million [27]. It was ranked 174th in the United Nations Human Development Index in 2017 [28]. The annual seasonal climate consists of a long dry season from November to May and a short rainy season between June and October. Sukuta is an area within the West Coast Region peri-urban conurbation, with a population of 47,048 in 2013, including 7,234 children aged under 5 years, and an average household size of 8.1 [27].

**Study design and sampling**

A cross-sectional, population-based study was conducted in Sukuta, in children aged <5 years, to determine the prevalence of scabies, pyoderma and fungal infections. Prior to the study, geospatial census data from 2013 was used to divide Sukuta into 37 geographical clusters of approximately 100 households (S1 Fig). In order to minimise selection bias, a one-stage, random cluster sampling method was used, sampling clusters in a randomly generated order. The latest census data (2013) on households was out of date, so was not used as a sampling list in each geographical cluster. Instead, all households in each selected cluster were approached for participation. Eligible participants were children <5 years of age sleeping in a compound within the geographical area of the cluster. Children <5 years were chosen as this age-group are below school age, and therefore more likely to be present during home visits. Participants were examined over a four-month period between May and September 2018. Sample size was determined by availability of study resources.

Information on socio-demographic factors, and possible risk factors for presence of skin infections were collected from participant’s parents prior to examination of the participant, and from the participant’s infant welfare card (if available). Socio-demographics recorded included age, sex, tribal group, mother’s educational level and household size. Possible risk factors and confounders recorded were breastfeeding status, birthweight (if known), household water source and distance, frequency of full body washing and ironing of clothes, whether clean clothes were worn every day, the presence in the compound of a handwashing area, an open fire for cooking, previous history of skin infections, burns, malnutrition or nutritional supplementation.

**Diagnosis, training and case definitions**

Study nurses underwent a 2-day training led by a physician, to introduce basic dermatology skills, recognise common paediatric skin complaints, and instruct on use of an adapted IMCI algorithm (S2 Fig) [29]. Participants were examined in the presence of their parents within the participant’s compound. This was often on private verandas, to ensure adequate natural light for skin examination, whilst maintaining privacy as far as possible. Clothes were removed where appropriate, though underwear was not removed unless necessary to closely examine or swab lesions. When examining sensitive areas or at the parents’ or participant’s request, examinations were performed indoors for more privacy. Scabies cases were diagnosed clinically...
based on the presence of pruritus and papules in a typical distribution. Pyoderma was defined as any skin lesion with evidence of pus or crusts. Infected scabies was diagnosed when scabies was present with evidence of inflammation and pyoderma in the same distribution. Fungal (dermatophyte) skin infection was diagnosed on the basis of the presence of round or oval flat scaly patches, with features typical of tinea infection. Where pyoderma was diagnosed, a wound swab was taken. All skin conditions identified were managed appropriately according to the algorithm and treatment guidelines (S2 Fig and S1 Table). Antibiotics were administered by study nurses in the field and follow-up of all cases was undertaken by the study clinician within one week.

Algorithm validity
To determine the sensitivity and specificity of the algorithm used, a subset of participants were selected opportunistically and re-examined by a general physician with experience of diagnosing paediatric dermatological conditions in The Gambia, blinded to the nurse’s diagnosis. A sample size of 123 was calculated to be sufficient to detect a sensitivity of 80% compared to the gold standard examination with a precision of 10% assuming a prevalence of 50% of each condition.

Swab collection and bacteriological methods
Wound swabs were taken to determine the presence of *S. aureus* and/or GAS by microbiological culture. A single swab was taken from the largest present pyoderma lesion; wounds were superficially cleaned with saline and crusts were lifted to swab the base. Samples were collected using Nylon flocked swabs (Copan) stored in liquid Amies transport medium. These were chosen over standard cotton swabs as future molecular/microbiome studies are planned. Swabs were transported in a cold box to the MRC Unit The Gambia at LSHTM (MRCG) for same-day culturing on 5% sheep’s blood agar and incubated at 37˚C overnight. Purity plates were done for mixed infection. Identification by catalase and confirmation for either *S. aureus* or GAS was done using the Remel Staphaurex Plus and Streptex latex agglutination tests, respectively. Antibiotic sensitivity pattern was obtained using disc diffusion according to CLSI methods [30] in line with the MRCG clinical diagnostic laboratory wound culture standard operating procedures. Only *S. aureus* and beta-haemolytic streptococci were considered relevant organisms to record.

Investigation of seasonality
Data collection spanned both the dry and rainy season (defined as after the 26th of June 2018 when the first rains of the year occurred), allowing for comparison between the prevalence, and the proportion of pyoderma caused by *S. aureus* and GAS between these two periods. In addition, the first cluster sampled was subsequently resampled at the end of the study in order to directly compare prevalence in the same population before and after the start of the rainy season.

To further investigate seasonal change in pyoderma in The Gambia, we also leveraged the presence of existing medical records from a rural primary healthcare clinic run by MRCG to perform a post-hoc analysis. The MRC Keneba clinic, in West Kiang, provides free primary care and has been collecting electronic medical records since 2009 [31]. Records on the number of presentations per month of <5 year olds for skin complaints (including impetigo, cutaneous abscess, furuncle or carbuncle, and cellulitis) between 2011 and 2018 were interrogated to ascertain if seasonal variation was observed.
Data collection and statistical analysis

A questionnaire was delivered to participant’s parents to collect information on socio-demographics and risk factors for skin diseases. Data were collected on tablet computers using RED-Cap electronic data capture tools hosted at MRCG [32]. Data were analysed using Stata version 15.1. The cluster random sampling method was corrected for using the `svyset` command. Adjusted binomial exact confidence intervals were calculated for prevalence estimates. Associations between socio-demographic and other risk factors and skin infections were investigated using multivariable logistic regression models. Variables and category levels were selected for inclusion using forwards and backwards stepwise regression, without `svyset` correction, using a likelihood ratio test significance level of $p<0.2$. The multivariable models were then rerun with `svyset` correction to obtain adjusted odds ratios (aOR). All socio-demographics and risk factors were categorical variables, except household size, which was continuous. The population attributable risk (PAR) [33] for scabies as a cause of pyoderma was calculated. The adjusted prevalence ratio (aPR) for the prevalence of skin conditions before and after the start of the rainy season were estimated using a multivariable Poisson regression model, adjusting for socio-demographics, except when comparing the first cluster before and after the rains, which was unadjusted. Predicted probabilities of the presence of skin conditions by week sampled were calculated using the `margins` command following Poisson regression models adjusting for socio-demographics. Sensitivity, specificity and kappa statistic were calculated to determine the accuracy of the diagnostic algorithm compared to the diagnosis reached by the physician. Significance was set at $p<0.05$.

Ethical considerations

Ethical approval for the study was provided by The Gambia Government/MRC Joint Ethics Committee (SCC1587). Written or thumb-printed informed consent was obtained in a local language from a parent for all participants included in the prevalence study prior to involvement.

Results

Prevalence of skin infections

A total of 1441 participants from 9 clusters were examined between May and September 2018. The mean number of participants per cluster was 160.1 (range 62–306). Overall, 47.9% of participants were male and the mean age was 28.9 (SD 16.7) months. Scabies prevalence was 15.9% (95% CI 12.2–20.4), pyoderma prevalence was 17.4% (95% CI 10.4–27.7), and fungal infection prevalence was 9.7% (95% CI 6.6–14.0). The prevalence of scabies, pyoderma and fungal infections by socio-demographic characteristics is presented in Table 1. Prevalence of co-infections are presented in S2 Table.

Associations with socio-demographic and other risk factors

Table 2 shows the aOR for skin infection presence by socio-demographic and other risk factors. The odds of scabies and fungal infections were significantly lower in females than males (aOR 0.70, 95% CI 0.61–0.82 and aOR 0.44, 95% CI 0.32–0.61 respectively). Pyoderma increased with age (aOR 3.13 of pyoderma in 3–4 years compared to <1 year, 95% CI 2.00–4.88), was higher in Serehule children compared to Mandinka children (aOR 1.99, 95% CI 1.16–3.41) and increased with household size (aOR 1.03, 95% CI 1.01–1.06). Various behavioural risk factors were identified, including lower odds of scabies in those whose clothes are always ironed (aOR 0.26, 95% CI 0.07–0.98), and higher odds of fungal infections in
those not wearing freshly washed clothes every day (aOR 13.5, 95% CI 3.22–26.60). Current breastfeeding was protective against fungal infections, but increased the odds of scabies (aOR 1.67, 95% CI 1.12–2.49). A history of previous skin infections increased the odds of all three infections. Results from the univariable regression models are presented in S3 Table. Results from the forwards and backwards stepwise regression procedures are presented in S5 and S6 Tables.

The presence of pyoderma was significantly associated with scabies infestation (aOR 2.74, 95% CI 1.61–4.67) as shown in Table 3. The PAR of scabies as a cause of pyoderma was 14.7% (95% CI 5.1–23.3).

**Pyoderma swab culture results**

Pyoderma swabs were taken from 250 participants (one participant diagnosed with pyoderma was not swabbed). Overall, either *S. aureus* or GAS was cultured from 90.0% of swabs, with 80.8% positive for *S. aureus*, and 50.8% for GAS (41.6% were positive for both). Additionally, two samples were positive for other beta-haemolytic streptococci (one group D streptococcus and one non-groupable *Streptococcus* species). The proportion of GAS and *S. aureus* causing pyoderma were similar regardless of scabies infection (Table 3). All GAS isolates were sensitive to penicillin, while 99% of *S. aureus* isolates were methicillin-sensitive. Antibiotic sensitivities of isolates are presented in S3 and S4 Figs. Swabs taken from the lower limbs were significantly more likely to be positive for GAS than swabs from other sites (S7 Table).
Table 2. Adjusted odds ratios for socio-demographic and other risk factors potentially associated with skin infections.

|                      | Scabies       |                  | Pyoderma     |                  | Fungal       |                  |
|----------------------|---------------|-----------------|--------------|-----------------|--------------|-----------------|
|                      | aOR | p value | 95% CIs | aOR | p value | 95% CIs | aOR | p value | 95% CIs |
| Sex                  |     |         |         | aOR | p value | 95% CIs | aOR | p value | 95% CIs |
| Male                 | ref | ref     | ref     | ref | ref     | ref     | ref | ref     | ref     |
| Female               | 0.70 | 0.001* | 0.61–0.82 | 1.22 | 0.201 | 0.88–1.70 | 0.44 | <0.001** | 0.32–0.61 |
| Age category         |     |         |         | aOR | p value | 95% CIs | aOR | p value | 95% CIs |
| <1 year              | ref | ref     | ref     | ref | ref     | ref     | ref | ref     | ref     |
| 1–2 years            | 0.91 | 0.634 | 0.60–1.40 | 1.98 | 0.086 | 0.89–4.43 | 0.48 | 0.006* | 0.30–0.76 |
| 2–3 years            | 0.81 | 0.529 | 0.38–1.72 | 2.85 | <0.001** | 1.90–4.26 | 0.55 | 0.026* | 0.33–0.91 |
| 3–4 years            | 0.87 | 0.670 | 0.42–1.79 | 3.13 | <0.001** | 2.00–4.88 | 0.79 | 0.509 | 0.36–1.74 |
| Tribe                |     |         |         | aOR | p value | 95% CIs | aOR | p value | 95% CIs |
| Mandinka             | ref | ref     | ref     | ref | ref     | ref     | ref | ref     | ref     |
| Serehule             | 1.72 | 0.044* | 1.02–2.90 | 1.99 | 0.018* | 1.16–3.41 | 0.39 | 0.105 | 0.12–1.28 |
| Other                | 0.42 | 0.020* | 0.21–0.84 | 0.39 | 0.105 | 0.12–1.28 | 0.74 | 0.107 | 0.05–1.30 |
| Mean household size  |     |         |         |     |         |         |     |         |         |
| ref                  | 1.03 | 0.014* | 1.01–1.06 | 1.03 | 0.014* | 1.01–1.06 | 1.03 | 0.014* | 1.01–1.06 |
| Mother’s education†  |     |         |         | aOR | p value | 95% CIs | aOR | p value | 95% CIs |
| None                 | ref | ref     | ref     | ref | ref     | ref     | ref | ref     | ref     |
| Arabic school only   | 0.60 | 0.007* | 0.43–0.83 | 0.48 | 0.040* | 0.24–0.96 | 0.69 | 0.126 | 0.42–1.14 |
| Secondary only       |     |         |         |     |         |         |     |         |         |
| Currently breastfeeding | No | ref     | ref     | ref | ref     | ref     | ref | ref     | ref     |
| Yes                  | 1.67 | 0.017* | 1.12–2.49 | 0.52 | 0.043* | 0.27–0.98 | 1.80 | 0.006* | 1.25–2.59 |
| Low birth weight (<2.5kg) |     |         |         |     |         |         |     |         |         |
| No                   | ref | ref     | ref     | ref | ref     | ref     | ref | ref     | ref     |
| Yes                  | 1.67 | 0.017* | 1.12–2.49 | 0.52 | 0.043* | 0.27–0.98 | 1.80 | 0.006* | 1.25–2.59 |
| Water source         |     |         |         |     |         |         |     |         |         |
| Tap                  | ref | ref     | ref     | ref | ref     | ref     | ref | ref     | ref     |
| Well                 | 1.41 | 0.434 | 0.54–3.73 | 1.41 | 0.434 | 0.54–3.73 | 1.41 | 0.434 | 0.54–3.73 |
| Clean clothes        |     |         |         |     |         |         |     |         |         |
| Every day            | ref | ref     | ref     | ref | ref     | ref     | ref | ref     | ref     |
| Not every day        | 13.50 | 0.003* | 3.22–56.60 | 13.50 | 0.003* | 3.22–56.60 | 13.50 | 0.003* | 3.22–56.60 |
| Clothes ironed       |     |         |         |     |         |         |     |         |         |
| Never                | ref | ref     | ref     | ref | ref     | ref     | ref | ref     | ref     |
| Always               | 0.26 | 0.048* | 0.07–0.98 | 0.26 | 0.048* | 0.07–0.98 | 0.26 | 0.048* | 0.07–0.98 |
| Handwashing area in compound |     |         |         |     |         |         |     |         |         |
| No                   | ref | ref     | ref     | ref | ref     | ref     | ref | ref     | ref     |
| Yes                  | 0.74 | 0.107 | 0.50–1.09 | 0.74 | 0.107 | 0.50–1.09 | 0.74 | 0.107 | 0.50–1.09 |
| Open fire in compound |     |         |         |     |         |         |     |         |         |
| No                   | ref | ref     | ref     | ref | ref     | ref     | ref | ref     | ref     |
| Yes                  | 1.51 | 0.021* | 1.09–2.10 | 1.40 | 0.018* | 1.08–1.82 | 1.51 | 0.021* | 1.09–2.10 |
| Previous skin infection |     |         |         | aOR | p value | 95% CIs | aOR | p value | 95% CIs |
| None                 | ref | ref     | ref     | ref | ref     | ref     | ref | ref     | ref     |
| One                  | 3.09 | <0.001** | 2.19–4.64 | 1.94 | 0.002* | 1.40–2.69 | 2.34 | 0.005* | 1.41–3.87 |
| More than one        | 4.51 | <0.001** | 2.63–7.73 | 2.34 | 0.002* | 1.53–3.57 | 2.14 | 0.072 | 0.92–5.02 |
| History of burn      |     |         |         |     |         |         |     |         |         |
| No                   | ref | ref     | ref     | ref | ref     | ref     | ref | ref     | ref     |
| Yes                  | 0.65 | 0.154 | 0.35–1.22 | 0.65 | 0.154 | 0.35–1.22 | 0.65 | 0.154 | 0.35–1.22 |
| History of malnutrition |     |         |         |     |         |         |     |         |         |
| No                   | ref | ref     | ref     | ref | ref     | ref     | ref | ref     | ref     |
| Yes                  | 0.67 | 0.243 | 0.32–1.40 | 0.67 | 0.243 | 0.32–1.40 | 0.67 | 0.243 | 0.32–1.40 |
| History of nutritional supplementation |     |         |         |     |         |         |     |         |         |
| No                   | ref | ref     | ref     | ref | ref     | ref     | ref | ref     | ref     |
| Yes                  | 0.32 | 0.170 | 0.06–1.83 | 0.32 | 0.170 | 0.06–1.83 | 0.32 | 0.170 | 0.06–1.83 |

Reported values are from multivariable logistic regression models, including only variables and category levels selected by stepwise regression with an elimination level of p>0.2. Grey cells represent variables not included in each multivariable analysis. For all three skin infections, forwards and backwards stepwise regression produced the same final model (S5 and S6 Tables). Sex and age group were included in all models, and correction for cluster sampling design was done on the final models. Data from univariable logistic regression models are shown in S3 Table. ref = reference category used; aOR = adjusted odds ratio

*significant at p<0.05
**significant at p<0.001
†English school attendance

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Seasonal variation of skin infection prevalence

The respective prevalence of scabies, pyoderma and fungal infection were 15.3%, 8.9% and 14.4% before the start of the rainy season, compared to 16.3%, 23.1% and 6.6% after. The prevalence of scabies did not change significantly (aPR 1.08, 95% CI 0.70–1.67), whereas pyoderma prevalence significantly increased following the start of the rains (aPR 2.42, 95% CI 1.39–4.23). Fungal infection prevalence fell significantly (aPR 0.44, 95% CI 0.32–0.60) (S8 Table). The PAR of scabies as a cause of pyoderma was 18.5% (95% CI 4.1–30.8) before the start of the rainy season compared to 13.6% (95% CI -0.7–25.8) afterwards. The prevalence of scabies, pyoderma and fungal skin infections by week sampled are presented in Fig 1. The increase in pyoderma prevalence during the rainy season was confirmed by resampling of the first cluster sampled (7.9% vs. 21.7%, PR 2.74, 95% CI 1.23–6.12). The start of the rains did not significantly affect the proportion of *S. aureus* or GAS detected (S8 Table).

The data obtained from the MRCG Keneba clinic records of presentations per month of children <5 for all skin complaints are presented in Fig 2. The number of presentations peaked each year between June and December, coinciding with the rainy season.

Sensitivity and specificity of the diagnostic algorithm

Diagnosis by nursing staff using the algorithm (S2 Fig) was 97.1% sensitive and 96.6% specific for the diagnosis of pyoderma, and 83.3% sensitive and 97.0% specific for non-infected scabies. Sensitivity and specificity for the pyoderma and scabies co-infection were 81.3% and 97.2% respectively. Sensitivity of detection of fungal infections was lower at 66.7%, but specificity was 95.8%. Full results including the kappa statistic for inter-rater agreement are presented in S9 Table.

Discussion

We found a high burden of skin infections among Gambian children and that pyoderma increased significantly during the rainy season. The scabies prevalence we observed of 15.9% is consistent with other studies from sSA [15, 16, 18–20], but the overall prevalence of pyoderma of 17.4% is higher than the estimated median impetigo prevalence of 7% for Africa [21]. Studies in other settings have found a strong association between scabies and pyoderma, with one study in Fiji finding a PAR of scabies as a cause of impetigo of 93.1% [10]. We observed a PAR of 14.7%, indicating that scabies may play a less significant role in pyoderma rates in this setting.
Fig 1. Predicted probabilities of skin infection prevalence by week examined.

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Fig 2. Number of skin complaint presentations in under 5s at MRCG Keneba clinic by month 2011–2018.

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setting. Interventions to reduce scabies, shown to be effective elsewhere [34–36], may still be worthwhile given the relatively high scabies prevalence.

The 8.9% pyoderma prevalence in children examined before the start of the rainy season was consistent with previous estimates for Africa [21]. However, in those examined during the rainy season, the prevalence rate was significantly higher (23.1%). This effect was confirmed in the cluster that was sampled both before and after the start of the rainy season. Furthermore, a strong seasonal trend was seen at the rural MRCG Keneba clinic in paediatric skin presentations between 2011 and 2018, which supports the hypothesis that bacterial skin infections are seasonal in The Gambia, a finding also observed in 1980 [17].

Very little microbiology data exist from pyoderma cases in Africa [3, 21]. By taking swabs from pyoderma lesions, we were able to provide robust estimates for the prevalence of S. aureus and GAS pyoderma. 50.8% of swabs were positive for GAS, which is lower than the global median of 74% noted previously from the limited heterogeneous studies available [21]. However, with the high pyoderma prevalence during the rainy season, this still indicates a significant exposure of children <5 to GAS and S. aureus annually. Better understanding the carriage and transmission of these organisms within households and communities in The Gambia is required to design preventative interventions to reduce the disease burden. Our antimicrobial susceptibility data showed reassuringly low rates of methicillin-resistant S. aureus which is consistent with other data we and others have recently published [37, 38]. Of greater concern was the resistance to erythromycin (13.3%) and co-trimoxazole (24.2%), especially as MDA of azithromycin is recommended by the WHO to eradicate trachoma and yaws [39, 40], and use may increase in African settings as evidence of impact on <5 mortality increases [41, 42]. As co-trimoxazole is widely used in The Gambia, the reduced susceptibility observed was unsurprising [43].

The reason for the seasonal increase in skin infections is not clear, but may reflect behavioural changes between seasons or environmental changes enhancing bacterial proliferation on skin. Indoor space can be limited within households, and during the rainy season people may spend more time in overcrowded indoor areas, increasing skin-to-skin contact. This could partly explain the observed increase, although scabies prevalence did not increase correspondingly as would be expected. Data were not collected on insect bites, but increased insect activity during the rainy season may play a more significant role than scabies in this setting. Further studies are required to explore the exact mechanism for this observation. We also observed associations between increased age, household size, history of previous infections, and certain tribal groups with higher pyoderma rates, some of which have been observed elsewhere [13, 44]. These suggest that individual’s behaviour may also be important in pyoderma transmission.

Additionally, associations between scabies and fungal infections and other risk factors were seen. Current breastfeeding increased the odds of scabies, which may reflect higher risk due to greater skin contact with mothers who may be the source of infection. Ironing of clothes reduced the odds of scabies, which could suggest that some transmission of mites may occur through clothes, in spite of the common understanding that scabies is transmitted only through direct contact. The largest effect-size was seen in participants not wearing freshly washed clothes every day, which increased the odds of fungal infections. These may represent potential targets for behavioural interventions to reduce these skin infections.

In low-resourced healthcare settings, nurses can be expected to diagnose and treat common conditions with little or no dermatology training [21, 29]. Nurses were used in our study for the diagnosis of the skin conditions rather than physicians to determine whether training in the use of a diagnostic algorithm for skin conditions could be effective in this setting. The sensitivity and specificity of the algorithm used were consistent with the validation study [29].
confirming that such algorithms for nurses can be effective diagnostic tools. Wider introduction and evaluation of such algorithms could be carried out in The Gambia and similar settings, with a view to increasing diagnosis and treatment, cost-effectiveness, and the effects on other GAS disease. An impact on reducing inappropriate antibiotic prescribing

For pyoderma in the absence of guidelines should also be considered, given the likely contribution to antimicrobial resistance [45].

Our study has several limitations. We sampled from one peri-urban population, so our findings may not be generalisable, particularly to rural settings, where the majority of Gambians live. Furthermore, as we sampled children <5 in this study due to practical constraints, we were unable to include older school-age children, in whom pyoderma prevalence may be higher [10, 22]. Older children would be an important consideration in future studies, especially as children <5 years would currently be excluded from mass drug administration (MDA) of ivermectin for scabies due to weight-based dosing rules [46]. This younger age group may, however, benefit from MDA with antibiotics such as azithromycin to reduce the burden of pyoderma [47, 48].

No up-to-date list of households or residents was available for the area, and it was not feasible to enumerate the area in advance of the study. We therefore used a random cluster sampling method, dividing Sukuta into clusters of approximately 100 households based on 2013 census data, without detailed within household population metrics. This reduced the precision of the prevalence confidence intervals and prevented analysis of results with respect to households. Accordingly, we also did not have accurate data on children missing during study visits, which had the potential to introduce some selection bias. The study was conducted over four months, so we were unable to determine whether the observed seasonal variation is cyclical on an annual basis or changes year-to-year. We were only able to resample one cluster to compare prevalence directly, and given that prevalence varied by cluster, we should interpret the results cautiously. Additionally, the associations observed between risk factors and skin infections should be interpreted with caution as question responses were open to recall bias, and there may have been residual confounding factors not measured. Finally, the subset used for the algorithm validation sub-study were selected opportunistically, which may have been open to selection bias. Despite these limitations, this study provides good baseline data for the prevalence of these common skin infections in a country where no other recent data exists and also provides valuable microbiological data on skin pathogens, which is lacking from LMIC [1–3].

This study has shown that skin infections are common in The Gambia, particularly pyoderma caused by *S. aureus* and GAS during the rainy season. Skin infections may be overlooked, particularly in LMIC where there are other pressing health concerns, but with such high prevalence of pyoderma, they may represent a crucial early exposure to potentially serious pathogens. The high incidence of invasive *S. aureus* infections in Gambian children is increasingly recognised and can be triggered by skin infections [38]. The exact mechanism by which RHD follow GAS infection is still not understood, but repeated GAS skin infections may be important [49]. It is plausible that exposure to GAS through the skin at a young age may be a step in the aetiology of RHD [8, 9], particularly as GAS pyoderma is known to trigger the better understood, immune-complex mediated APSGN, which is also epidemiologically linked to scabies [23, 50].

**Conclusions**

Treating skin infections for their own sake is justification enough, but since GAS and *S. aureus* pyoderma can lead to serious disease, diagnosing and treating them as an upstream target to
prevent invasive infections and non-suppurative complications may also be a cost-effective way to prevent these more serious conditions. Increased research into GAS carriage, transmission dynamics, and correlates of protection from disease is therefore warranted in African settings.

Supporting information

S1 Checklist. STROBE checklist.
(DOCX)

S1 Fig. The Sukuta area divided into 37 geographical clusters with approximately equal number of compounds from 2013 geospatial census data, used for one-stage cluster sampling method.
(TIF)

S2 Fig. Diagnostic algorithm for common skin conditions adapted from IMCI algorithm designed and validated by Steer et al. (2009).
(TIF)

S3 Fig. *Staphylococcus aureus* isolate antibiotic sensitivities.
(TIF)

S4 Fig. Group A streptococcus isolate antibiotic sensitivities.
(TIF)

S1 Table. Treatment guidelines used for common skin conditions.
(DOCX)

S2 Table. Prevalence of infected scabies, and co-infection with scabies, pyoderma and fungal infections.
(DOCX)

S3 Table. Odds ratios for socio-demographic and other risk factors potentially associated with skin infections in univariable logistic regression models.
(DOCX)

S4 Table. Adjusted odds ratios for socio-demographic and other risk factors potentially associated with skin infections in global multivariable logistic regression models including all variables.
(DOCX)

S5 Table. Forwards stepwise logistic regression steps.
(DOCX)

S6 Table. Backwards elimination stepwise logistic regression steps.
(DOCX)

S7 Table. Swab positivity by swab site.
(DOCX)

S8 Table. Prevalence rate ratios for presence of skin infections by whether examined before or after the start of the rainy season using Poisson regression.
(DOCX)

S9 Table. Sensitivity, specificity and kappa statistic for each skin diagnosis made by trained nurses using the adapted skin condition diagnostic algorithm as compared with the
diagnosis of a physician.

S1 Dataset. Main study dataset.

S2 Dataset. Skin complaint presentation dataset from MRCG Keneba clinic.

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References

1. Zuhlke LJ, Beaton A, Engel ME, Hugo-Hamman CT, Karthikeyan G, Katzenellenbogen JM, et al. Group A Streptococcus, Acute Rheumatic Fever and Rheumatic Heart Disease: Epidemiology and Clinical Considerations. Curr Treat Options Cardiovasc Med. 2017; 19(2):15. Epub 2017/03/13. https://doi.org/10.1007/s11936-017-0513-y PMID: 28285457; PubMed Central PMCID: PMC5346434.
20. Walker SL, Lebas E, De Sario V, Deyasso Z, Doni SN, Marks M, et al. The prevalence and association with health-related quality of life of tungiasis and scabies in schoolchildren in southern Ethiopia. PLoS Negl Trop Dis. 2017; 11(8):e0005808. Epub 2017/08/05. https://doi.org/10.1371/journal.pntd.0005808 PMID: 28771469; PubMed Central PMCID: PMC5557602.

21. Bowan AC, Mahe A, Hay RJ, Andrews RM, Steer AC, Tong SY, et al. The Global Epidemiology of Impetigo: A Systematic Review of the Population Prevalence of Impetigo and Pyoderma. PLoS One. 2015; 10(8):e0136789. Epub 2015/09/01. https://doi.org/10.1371/journal.pone.0136789 PMID: 26317533; PubMed Central PMCID: PMC4552802.

22. Romani L, Steer AC, Whitfield MJ, Kaldor JM. Prevalence of scabies and impetigo worldwide: a systematic review. Lancet Infect Dis. 2015; 15(8):960–7. Epub 2015/06/20. https://doi.org/10.1016/S1473-3099(15)00132-2 PMID: 26088526.

23. Walker MJ, Barnett TC, McArthur JD, Cole JN, Gillen CM, Henningham A, et al. Disease manifestations and pathogenic mechanisms of Group A Streptococcus. Clin Microbiol Rev. 2014; 27(2):264–301. Epub 2014/04/04. https://doi.org/10.1128/CMR.00101-13 PMID: 24696436; PubMed Central PMCID: PMC3993104.

24. Steer AC, Law I, Matatolu L, Beall BW, Carapetis JR. Global emm type distribution of group A streptococci: systematic review and implications for vaccine development. Lancet Infect Dis. 2009; 9(10):611–6. Epub 2009/08/26. https://doi.org/10.1016/S1473-3099(09)70179-1 PMID: 19778673.

25. Barth DD, Engel ME, Whitelaw A, Abdissa A, Sadoh WE, Ali SK, et al. Rationale and design of the AfriStrep study. BMJ Open. 2013; 2(6):e010248. Epub 2013/02/27. https://doi.org/10.1136/bmjopen-2013-010248 PMID: 23916994; PubMed Central PMCID: PMC4769387.

26. Coulibaly O, L’Ollivier C, Piarroux R, Ranque S. Epidemiology of human dermatophytoses in Africa. Medical mycology. 2016; 56(2):145–61. Epub 2017/10/11. https://doi.org/10.1093/mmy/myx046 PMID: 28992062.

27. GBoS. The Gambia 2013 Population and Housing Census. Kanifing: The Gambia Bureau of Statistics; 2013.

28. The Human Development Reports Office. Human Development Reports [Online]. New York: UNDP; 2018 [cited 2019 04 Jan]. Available from: http://hdr.undp.org/en/composite/HDI.

29. Steer A. Validation of an Integrated Management of Childhood Illness algorithm for managing common skin conditions in Fiji. Bulletin of the World Health Organization. 2009; 87(3):173–9. https://doi.org/10.2471/BLT.08.052712 PMID: 19377712.

30. CLSI. M02 Performance Standards for Antimicrobial Disk Susceptibility Tests. Wayne, PA: Clinical and Laboratory Standards Institute, 2018.

31. Hennig BJ, Unger SA, Dondeh BL, Hassan J, Hawkesworth S, Jarjou L, et al. Cohort Profile: The Kiang West Longitudinal Population Study (KWLPS)-a platform for integrated research and health care provision in rural Gambia. International journal of epidemiology. 2017; 46(2):e13. Epub 2015/11/13.

32. Harris PA, Taylor R, Thielke R, Payne J, Gonzalez N, Conde JG. Research electronic data capture [REDCap]—a metadata-driven methodology and workflow process for providing translational research informatics support. J Biomed Inform. 2009; 42(2):377–81. Epub 2008/10/22. https://doi.org/10.1016/j.jbi.2008.08.010 PMID: 18929886; PubMed Central PMCID: PMC2700030.

33. Northridge ME. Public health methods—attributable risk as a link between causality and public health action. Am J Public Health. 1995; 85(9):1202–4. Epub 1995/09/01. https://doi.org/10.2105/ajph.85.9.1202 PMID: 7661224; PubMed Central PMCID: PMC1615585.

34. Marks M, Taotao-Wini B, Satorara L, Engelmann D, Nasi T, Mabey DC, et al. Long Term Control of Scabies Fifteen Years after an Intensive Treatment Programme. PLoS Negl Trop Dis. 2015; 9(12):e0004246. Epub 2015/12/02. https://doi.org/10.1371/journal.pntd.0004246 PMID: 26624616; PubMed Central PMCID: PMC4664486.

35. Romani L, Whitefield MJ, Korovueta J, Kama M, Wand H, Tikoduadua L, et al. Mass Drug Administration for Scabies Control in a Population with Endemic Disease. N Engl J Med. 2015; 373(24):2305–13. Epub 2015/12/10. https://doi.org/10.1056/NEJMoa1500987 PMID: 26650152.

36. Marks M, Toloka H, Baker C, Kositz C, Asugeni J, Puiahi E, et al. Randomised trial of community treatment with azithromycin and ivermectin mass drug administration for control of scabies and impetigo. Clin Infect Dis. 2018. Epub 2018/07/10. https://doi.org/10.1093/cid/ciy574 PMID: 29985978.

37. Darboe S, Dobrzeniecki S, Jarju S, Jallow M, Mohammed NI, Watthuo M, et al. Prevalence of Panton-Valentine Leukocidin (PVL) and Antimicrobial Resistance in Community-Acquired Clinical Staphylococcus aureus in an Urban Gambian Hospital: A 11-Year Period Retrospective Pilot Study. Front Cell Infect.
38. Odutola A, Bottomley C, Zaman SA, Lindsay J, Shah M, Hossain I, et al. Staphylococcus aureus Bacteremia in Children of Rural Areas of The Gambia, 2008–2015. Emerg Infect Dis. 2019; 25(4):701–9. Epub 2019/03/19. https://doi.org/10.3201/eid2504.180935 PMID: 30882307.

39. WHO. Trachoma Fact Sheet: World Health Organization; 2019 [cited 2019 14 Aug]. Available from: https://www.who.int/news-room/fact-sheets/detail/trachoma.

40. WHO. Eradication of yaws—the Morges strategy. Releve epidemiologique hebdomadaire. 2012; 87(20):189–94. Epub 2012/05/18. PMID: 24349800.

41. Keenan JD, Bailey RL, West SK, Arzika AM, Hart J, Weaver J, et al. Azithromycin to Reduce Childhood Mortality in Sub-Saharan Africa. N Engl J Med. 2018; 378(17):1583–92. Epub 2018/04/26. https://doi.org/10.1056/NEJMoA1715474 PMID: 29694816; PubMed Central PMCID: PMC5849140.

42. Tam CC, Offeddu V, Lim JM, Voo TC. One drug to treat them all: ethical implications of the MORDOR trial of mass antibiotic administration to reduce child mortality. J Glob Health. 2019; 9(1):010305. Epub 2019/01/16. https://doi.org/10.7189/jogh.09.010305 PMID: 30643634; PubMed Central PMCID: PMC6318831 form at www.icmje.org/coi_disclosure.pdf (available on request from the corresponding author) and declare no conflict of interest).

43. Tadesse BT, Ashley EA, Ongarello S, Havumaki J, Wijegoonewardena M, Gonzalez LJ, et al. Antimicrobial resistance in Africa: a systematic review. BMC Infect Dis. 2017; 17(1):616. Epub 2017/09/13. https://doi.org/10.1186/s12879-017-2713-1 PMID: 28893183; PubMed Central PMCID: PMC5594539.

44. Bowen AC, Tong SY, Chatfield MD, Carapetis JR. The microbiology of impetigo in indigenous children: associations between Streptococcus pyogenes, Staphylococcus aureus, scabies, and nasal carriage. BMC Infect Dis. 2014; 14:727. Epub 2015/01/01. https://doi.org/10.1186/s12879-014-0727-5 PMID: 25551178; PubMed Central PMCID: PMC4299569.

45. Tong SY, Varrone L, Chatfield MD, Beaman M, Giffard PM. Progressive increase in community-associated methicillin-resistant Staphylococcus aureus in Indigenous populations in northern Australia from 1993 to 2012. Epidemiol Infect. 2015; 143(7):1519–23. Epub 2014/10/11. https://doi.org/10.1017/S0950268814002611 PMID: 25302939.

46. Wilkins AL, Steer AC, Cranwicks N, Gwee A. Question 1: Is it safe to use ivermectin in children less than five years of age and weighing less than 15 kg? Arch Dis Child. 2018; 103(5):514–9. Epub 2018/02/22. https://doi.org/10.1136/archdischild-2017-314505 PMID: 29463522.

47. Oluwalana C, Camara B, Bottomley C, Goodier S, Bojang A, Kampmann B, et al. Azithromycin in Labor Lowers Clinical Infections in Mothers and Newborns: A Double-Blind Trial. Pediatrics. 2017; 139(2). Epub 2017/01/29. https://doi.org/10.1542/peds.2016-2281 PMID: 28130432.

48. Fry AM, Jha HC, Lietman TM, Chaudhary JS, Bhatta RC, Elliott J, et al. Adverse and beneficial secondary effects of mass treatment with azithromycin to eliminate blindness due to trachoma in Nepal. Clin Infect Dis. 2002; 35(4):395–402. Epub 2002/07/30. https://doi.org/10.1086/341414 PMID: 12145722.

49. Cunningham MW. Streptococcus and rheumatic fever. Curr Opin Rheumatol. 2012; 24(4):408–16. Epub 2012/05/24. https://doi.org/10.1097/BOR.0b013e32835461d3 PMID: 22617826; PubMed Central PMCID: PMC3645882.

50. Svartman M, Finklea JF, Earle DP, Potter EV, Poon-King T. Epidemic scabies and acute glomerulonephritis in Trinidad. Lancet. 1972; 1(7744):249–51. Epub 1972/01/29. https://doi.org/10.1016/s0140-6736(72)90634-4 PMID: 4109709.