Surveillance in easy to access population subgroups as a tool for evaluating malaria control progress: A systematic review

Sanie S. S. Sesay, Emanuele Giorgi, Peter J. Diggle, David Schellenberg, David G. Lalloo, Dianne J. Terlouw

1 Malawi-Liverpool-Wellcome Trust Clinical Research Programme, Blantyre, Malawi, 2 Liverpool School of Tropical Medicine, Liverpool, United Kingdom, 3 Medical School, Lancaster University, Lancaster, United Kingdom, 4 Institute of Infection and Global Health, University of Liverpool, Liverpool, United Kingdom, 5 London School of Hygiene and Tropical Medicine, London, United Kingdom

* sanie.sesay@sanofi.com

Abstract

Background

The need for surveillance systems generating targeted, data-driven, responsive control efforts to accelerate and sustain malaria transmission reduction has been emphasized by programme managers, policy makers and scientists. Surveillance using easy-to-access population subgroups (EAGs) may result in considerable cost saving compared to household surveys as the identification and selection of individuals to be surveyed is simplified, fewer personnel are needed, and logistics are simpler. We reviewed available literature on the validation of estimates of key indicators of malaria control progress derived from EAGs, and describe the options to deal with the context specific bias that may occur.

Methods

A literature search was conducted of all documents reporting validation of estimates of malaria control indicators from EAG surveys before the 31st of December 2016. Additional records were identified through cross-reference from selected records, other applicable policy documents and grey literature. After removal of duplicates, 13, 180 abstracts were evaluated and 2,653 eligible abstracts were identified mentioning surveillance in EAGs, of which 29 full text articles were selected for detailed review. The nine articles selected for systematic review compared estimates from health facility and school surveys with those of a contemporaneous sample of the same population in the same geographic area.

Results

Review of the available literature on EAGs suitable for surveillance of malaria control progress revealed that little effort has been made to explore the potential approach and settings for use of EAGs; and that there was wide variation in the precision of estimates of control progress between and within studies, particularly for estimates of control intervention coverage. Only one of the studies evaluated the geospatial representativeness of EAG samples, or carried out geospatial analyses to assess or control for lack of geospatial representativeness.
Two studies attempted to measure the degree of bias or improve the precision of estimates by controlling for bias in a multivariate analysis; and this was only successful in one study. The observed variability in accuracy of estimates is likely to be caused by selection and/or information bias due to the inherent nature of EAGs. The reviewed studies provided insight into the design and analytical approaches that could be used to limit bias.

Conclusion

The utility EAGs for routine surveillance of progress in malaria control at the district or sub-district programmatic level will be driven by several factors including whether serial point estimates to measure transmission reduction or more precise geospatial distribution to track ‘hot-spots’ is required, the acceptable degree of precision, the target population, and the resources available for surveillance. The opportunities offered by novel geostatistical analyses and hybrid sampling frames to overcome bias justify a renewed exploration of use of EAGs for malaria monitoring and evaluation.

Background

The need for surveillance systems that inform accelerated and sustained control efforts to accelerate and sustain malaria transmission reduction has been emphasized by programme managers, policy makers and scientists. A key element of these surveillance systems will be their cost and whether they can easily be integrated with current malaria control activities. Routine health facility-based passive case reporting, for example through Health Management Information Systems (HMISs), has been and continues to be at the forefront of malaria surveillance [1, 2]. A well-functioning HMIS will provide regular data from all health facilities nationally allowing accurate measurement of malaria control progress across the healthcare system. This has largely not been the case for most HMISs in malaria endemic countries, with problems like incomplete reporting and lack of diagnostic confirmation being comparatively common [3, 4]. Malaria indicator surveys (MISs) provide single cross-sectional national assessments of disease burden [5], but are usually expensive and logistically demanding to undertake. The goal of MISs is to generate nationally representative estimates and are thus not powered to detect local-level variability[6, 7]. The interval between serial MISs also affect their direct relevance for monitoring short- and medium-term trends in malaria control progress. Supplementary approaches are thus needed to provide timely estimates of malaria control progress at the district and sub-district level, complementary to current malaria surveillance systems, particularly as malaria transmission intensity falls and its distribution becomes more localized [8].

Representative subsets of the population or disease at-risk groups routinely assemble at easily accessible locations (e.g. schools or health facilities) making them logistically attractive to sample [9]. Alternatively, representative subgroups or the whole population of interest may be easily accessible during public health intervention activities such as mass drug administration and catch-up vaccination campaigns [10]. The opportunistic nature of surveillance in the so-called Easy Access Groups (EAGs) could thus save costs by simplifying the identification and selection of individuals to be surveyed, requiring simplified logistics and fewer study personnel compared to household surveys [9, 11]. Evidence from school surveys indicate that EAGs are suitable for surveillance when they are potentially representative of an at-risk stratum of the
population [9]. However, there are concerns about the inherent biased nature of such a sample, as such non-probability samples depend on natural systems of selection which are likely to result in the selection of a non-representative sample of the population of interest [12]. In this systematic review, we studied the available literature on the validation of estimates of key indicators of malaria control progress [13] derived from EAGs, focusing on EAGs that may be suitable for surveillance at the district or (sub)district level.

Methods

Search strategy

We searched EMBASE® (EMBASE, Medline, EMBASE Classic), PubMed® and ScienceDirect® bibliographic databases without language restrictions from inception to 31st December 2016 for articles with the following search terms in their keywords, title or abstract: "malaria" AND "survey"; or "malaria" AND "surveillance", or "malaria" AND "monitoring" AND "evaluation"; or "malaria" AND "transmission" AND "measurement. We also searched the online WHO document centre [14] for relevant policy documents and for grey literature from the WHO historical documents database on malaria (1947–2000) [15]. We also included pertinent articles that were not picked up by our search from other sources including recommendation from key experts in the field of malaria surveillance.

We compiled the results into a searchable database in EndNote X8.0.1 (Thomson Reuters). We searched this database for abstracts detailing validation of estimates from EAGs predetermined to be most suitable for routine malaria surveillance at the (sub)district-level by a review of historical evidence of previous use for malaria surveillance. We also added EAGs that had not been previously used for malaria surveillance but demonstrated this potential through surveillance of other diseases. Selected EAGs were further validated by examination against general criteria used to evaluate the suitability of a surveillance system [16], adapted to malaria surveillance (Table 1). Based on our review we postulated that the following EAGs were suitable for the routine surveillance of malaria control progress (Table 2):

Table 1. Criteria evaluating the suitability of EAGs for malaria surveillance.

| Attribute                | Definition                                                                 |
|--------------------------|---------------------------------------------------------------------------|
| Usefulness               | Contributes to understanding the epidemiology of malaria in the study area. Generates a suitable prompt public health response by impacting policies and/or control response. |
| Cost-effective           | The direct and indirect costs should be justifiable in relation to the benefits attained. |
| Quality                  |                                                                           |
| Sensitivity              | The ability of the surveillance system to measure presence of relevant impact indicators. |
| Specificity              | The ability of the surveillance system to identify the absence of relevant impact indicators. |
| Representativeness       | Accurately reflects the spatio-temporal distribution of key health events and uptake of public health control measures in the population or key at-risk groups. |
| Timeliness               | Ability to provide timely estimates of key health events to guide control efforts. |
| Simplicity               | Easy to understand and implement.                                         |
| Flexibility              | Ability to be easily adapted to include new or emerging problems, other health events, population sub-groups or key disease at-risk groups. |
| Acceptability            | Willingness of persons conducting surveillance and those providing data to generate accurate, consistent and timely data. Acceptability to other key stakeholders, the community, health planners, donors, etc. |

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Table 2. Advantages and disadvantages of EAGs suitable for malaria surveillance.

| EAG                                      | Advantages                                                                 | Disadvantages                                                                 |
|------------------------------------------|----------------------------------------------------------------------------|-------------------------------------------------------------------------------|
| School children                          | • Age range of primary school children in Africa of 5 to 14 years captures the PfPR peak [84, 85]  
• Allows direct measurement of impact of malaria control interventions targeted at school children [86]  
• Extensively assessed historically [17, 87] and at the district and sub-district level [11, 88] | Substantial variations in primary school enrolment rates between different regions in sub-Saharan Africa [86] |
| Health facility attendees                |                                                                             |                                                                               |
| All health facility attendees            | • Less susceptible to problems of HMISs such as incomplete reporting and lack of diagnostic confirmation [3, 89] | • Representativeness of data on control progress from health facilities surveys will depend largely on health facility utilization rates [52, 90, 91] |
| Children coming for sick or "well" child visits | • Mostly infants which are a sensitive group to measure malaria transmission [92]  
• Can be used to directly assess coverage where immunization clinics have been used to distribute malaria control interventions [93] | • Blood sampling is required may have ethical considerations and may cause poor acceptance especially in children coming for well child visits  
• Same considerations for representativeness as above |
| Women attending ANC or coming for delivery | • Pregnant women are more susceptible to malaria regardless of endemicity making them a sensitive group to measure malaria transmission [19, 94]  
• Parity specific susceptibility suggest primigravidae are an even more sensitive at-risk sub-group [95–97]  
• ANC attendance is high and most women attend ANC at least once during their pregnancy [57]  
• PfPR at the first antenatal booking is likely to reflect population transmission pressure as these women are yet to receive control interventions targeted at malaria in pregnancy [98]  
• Blood sampling requirement at first ANC visit and at delivery can be used to assess PfPR and APR | • No integrated strategic approach to surveillance of malaria control in pregnancy currently so indicators need to be validated [99]  
• Relationship between the prevalence of peripheral and placental parasitaemia in pregnant women and that of the population is poorly understood [100]  
• Women with lower SES in developing countries are less likely to deliver in health facilities and this affects representativeness [101] |
| Population targeted by public health intervention/campaign | • Most of the population or at-risk group is available for sampling  
• Mass ITN distribution, national immunization days (NIDs), mass drug administration (MDA) and surveys for NTDs offer excellent opportunities to integrate malaria surveillance, and has been assessed with MDA for filariasis [20] and surveys for trachoma [102] | • Unlikely to be a source of continuous data |
| Population attending rural community markets | • Rural markets in large, centrally place towns offer an opportunity to survey a large potentially representative sample of the adult community of the surrounding area involving all social strata, and has not been assessed for malaria surveillance but in other diseases [103][42][104] | • Needs to be validated for malaria surveillance, and in urban settings |

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1. School children
2. Health facility attendees, including:
   a. All health facility attendees including accompanying persons
   b. Children coming for sick or routine "well" child visits
   c. Women attending ANC or coming for delivery
3. Population targeted by public health intervention campaign such as mass drug administration
4. Population attending rural community markets
We then searched the EndNote database for articles with the following keywords in their abstract:

1. "school" AND "survey", "school AND "surveillance", "school" AND "monitoring" AND "evaluation", and "school" AND "transmission" AND "measurement"

2. "health" AND "facility" OR "centre" AND "survey", health" AND "facility" OR "centre" AND "surveillance", "health" AND "facility" OR "centre" AND "monitoring" AND "evaluation", and "health" AND "facility" OR "centre" AND "transmission" AND "measurement"

3. "antenatal clinic" AND "survey", "antenatal clinic" AND "surveillance", "antenatal clinic" AND "monitoring" AND "evaluation", "antenatal clinic" AND "transmission" AND "measurement", "pregnancy" OR "delivery" AND "survey", "pregnancy" OR "delivery" AND "surveillance", "pregnancy" OR "delivery" AND "monitoring" AND "evaluation", and "pregnancy" OR "delivery" AND "transmission" AND "measurement"

4. "market" AND "survey", "market" AND "surveillance", "market" AND "monitoring" AND "evaluation", and "market" AND "transmission" AND "measurement"

5. "public health" AND "intervention" OR "campaign" AND "survey", "public health" AND "intervention" OR "campaign" AND "surveillance", "public health" AND "intervention" OR "campaign" AND "monitoring" AND "evaluation", and "public health" AND "intervention" OR "campaign" AND "transmission" AND "measurement"

**Inclusion criteria**

A total of 13,180 records were compiled into a searchable database, at which the key word search resulted in the selection of 2,653 eligible abstracts for further review. These abstracts were reviewed for specific mention of the comparison of malaria indicator estimates from an EAG sample with population sample (Fig 1) and 29 articles were selected for full text review.

**Exclusion criteria**

We searched for the full text of the selected 29 publications, and excluded studies in which estimates of malaria control indicators from EAGs were not compared to a contemporaneous random population sample from the same geographic area. Since the distribution of *Plasmodium falciparum* infection in the population is determined by environmental factors that influence the density of competent anopheline mosquitoes, location-specific vector behaviour, and human factors like at-risk status (e.g. age and pregnancy) and behaviour (e.g. ITN use) that increase exposure to infectious mosquito bites [17–19]; to increase the accuracy of EAG *Pf*PR estimates, we excluded all studies that did not compare EAG samples to population samples from the same age or other at-risk stratum.

**Selection of studies**

Twenty of the twenty-nine studies selected for full review satisfied one or more exclusion criteria and were not included in the systematic review (Fig 1). Six of the studies were excluded because the data collected was not sufficiently synchronous between the EAG and population sample [20–25]. In fourteen studies the validity of EAG estimates could not be determined either because the population sample was not random [26], the same indicators were not collected from both samples [27–31], or both samples were otherwise not comparable [32–39]. In the nine selected studies, information was recorded on the type of EAG, comparator...
\[n^1 = \text{EMBASE}(\text{EMBASE, Medline, EMBASE Classic}) + \text{PubMed} + \text{ScienceDirect}\]

\[n^2 = \text{WHO policy documents} + \text{WHO historical documents database} + \text{other sources}\]

\[n^3 = \text{School surveys} + \text{Health facility surveys} + \text{ANC and/or delivery surveys} + \text{Public health campaign surveys} + \text{Community market surveys}\]

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**Fig 1.** PRISMA flow diagram for studies comparing estimates between EAG and population surveys.

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In six studies, the data were not synchronous. In fourteen studies, the EAG estimates could not be validated because:

- the population sample was not random (1),
- similar indicators were not collected from both samples (5), or
- otherwise not possible to compare EAG and population (8).
population, sampling frame, sampling methodology, sample size and sampling units. Data on the first author, year of survey, survey site, year of publication, malaria transmission intensity (e.g. PfPR), and estimates of control progress were extracted for the systematic review.

Definitions

Anaemia prevalence rate (APR)–Proportion of the population with a haemoglobin measurement of <8 g/dL.

Antibody prevalence rate (AbPR)–Proportion of the population seropositive to defined malaria antigens.

Household bed net ownership–Proportion of households with at least one bed net.

Household ITN ownership–Proportion of households with at least one ITN.

Individual bed net use–Proportion of population that slept under a bed net the previous night.

Individual insecticide treated bed net (ITN) use–Proportion of population that slept under an ITN the previous night.

Indoor residual spraying (IRS) coverage–Proportion of households sprayed with IRS in the past 12 months.

Plasmodium falciparum prevalence rate (PfPR)–Proportion of the population with malaria infection detected by rapid diagnostic test (RDT), microscopy or polymerase chain reaction (PCR).

Sick child visit–Health facility visits during childhood for an illness episode.

Well child visit–Routine health facility visits that occur during childhood that may include immunizations, growth and development assessments, physical examination and other health risk assessments.

Statistical analysis

Data analysis was done using Stata version 13.1 (StataCorp, Texas, USA). Using the presented data from tables in the selected publications, we calculated point estimates of control progress indicators derived from EAGs and compared that to estimates from contemporaneous population samples. In one publication [40], due to absence of the numerator, we derived the numerator from the reported rates and the denominator, and then calculated point estimates and corresponding 95% confidence intervals. Where surveys were repeated either seasonally or after a specific period [41, 42], we presented these estimates separately to account for seasonal or temporal effect respectively. We assessed the degree of accuracy in estimates derived from EAG samples by examining the absolute difference in prevalence difference and corresponding 95% confidence intervals and Pearson’s $\chi^2$ p values. Mean prevalence was derived for the overall individual level estimates from the EAG and population samples. The estimates for PfPR were derived individually for each method of detection of parasitaemia e.g. blood film, rapid diagnostic test. Malaria endemicity was classified according to the revised Global Malaria Eradication Program classification [43]. Due to the inherent differences in EAGs and the paucity of studies, we did not derive pooled estimate effects for each malaria control indicator. To evaluate the effect of population coverage of control interventions and transmission intensity on the validity of EAG estimates of control interventions and PfPR respectively, where possible, we correlated the prevalence difference with the population prevalence. We also evaluated the potential for any of the EAG samples to misclassify an area into the wrong malaria endemicity category by comparing the classification of each area by population PfPR to that from EAG estimates.
Results

Description of studies

Nine studies were included in the systematic review (Table 3), all of which were from sites with intense stable or moderate stable malaria transmission. Six studies assessed the accuracy of estimates from health facilities [40–42, 44–46], two studies assessed the accuracy of estimates from school surveys [47, 48], and one study assessed the accuracy of estimates from antenatal clinics [49]. Three studies compared estimates from children less than 5 years old [40, 42, 44], two studies compared estimates from older children [47, 48], and four studies compared estimates from all presenting individuals at health facilities (including ANC) regardless of age [41, 45, 46, 49].

Comparison of estimates

Estimates of coverage of control interventions. Seven studies assessed the accuracy of estimates of coverage of control interventions. (Table 4) [40–42, 44, 47–49]. The estimates of coverage of different control interventions derived from EAGs were significantly higher than those of the population in three studies [40, 44, 47], except for the estimates of household ITN ownership which was concordant with the population in one of these studies [47]. In three studies, estimates of control intervention coverage were significantly lower in EAGs [41, 48, 49]. In one study, estimates derived from parents/guardians of children aged 6–30 months coming for well child visits in Malawi were concordant in the first year of survey (2005) but significantly higher in the second survey (2008) [42]. In 2005, the estimates of individual bed net use derived from this EAG (PR = 41.0%, 95% CI 38.9%, 47.4%) were slightly lower than that in the same age stratum in the population (PR = 45.4%, 95% CI 39.0%, 51.7%, p = 0.0339), though this difference is not significant due to overlapping confidence intervals. Similarly, the estimate of individual ITN use derived from the EAG in the same survey (PR = 36.7%, 95% CI 31.1%, 42.4%) was not significantly different from that of the population (PR = 41.0%, 95% CI 34.1%, 40.5%, p = 0.0311). The study by Stevenson et al [48] investigated the concordance in school and catchment area-based estimates of control intervention coverage across a range of circumferential distances around each school. Estimates of individual bed net use derived from school children living 601–1000m (PR = 31.3%, 95% CI 33.5%) and >1000m (PR = 33.4%, 95% CI 35.6%) from the school were not significantly different from those from school children within 600m of the school (PR = 33.4%, 95% CI 31.2%, 35.6%), indicating that inaccuracy remained relatively constant with changes in circumferential area within the school’s catchment area. In the same study, estimates of IRS coverage from school children living 601–1000m (PR = 70.7%, 95% CI 68.5%, 72.8%) and >1000m (PR = 72.9%, 95% CI 68.5%, 72.8%) from the school were not significantly different from those from school children within 600m (PR = 68.3%, 95% CI 66.1%, 70.4%) of the school again indicating the inaccuracy was not affected by circumferential area within the school’s catchment area.

Estimates of malaria morbidity. Six studies assessed the accuracy of estimates of malaria morbidity (Table 5) [41, 42, 44–46, 48]. All six studies evaluated estimates of Plasmodium falciparum prevalence rate (PfPR) either by rapid diagnostic test (RDT), microscopy or polymerase chain reaction (PCR). In the studies where PfPR was determined by microscopy, slides were double read [41, 44, 45] or single read by an expert microscopist [42]. As an additional measure, in two studies there was external quality control [41, 42], and in one study PCR was used to complement missing second reads and to disambiguate discordant species read results [45]. In three studies, estimates of PfPR derived from EAGs were significantly higher than those of the population [45, 46, 48]. In one study [44], estimates of PfPR derived from children
### Table 3. Description of studies comparing estimates between EAG and population surveys.

| Study | Year(s) | Country          | Geographic unit comparison | Malaria endemicity | EAG Participants | Population Participants | Sampling methods | Sampling Units, No. sampled |
|-------|---------|------------------|-----------------------------|-------------------|-------------------|--------------------------|------------------|----------------------------|
| Briand et al. [49] | 2014 | Laos Region Salavan | Moderate stable | Pregnant women living in households | 30 villages: n = 205 | Random selection of pregnant women invited to participate | Sampled, n = 545 |
| Gahutu et al. [44] | 2011 | Rwanda Sub-district Butare | Moderate stable | Children < 5 years living in households | Successive | 1 hospital: n = 101; 1 health centre: n = 103 |
| Hetzel et al. [45] | 2008/9 | Papua New Guinea Momase and Highlands | Moderate stable | All individuals living in households older than 5 months | Successive | 3 health centres: n = 304 |
| Karyana et al. [46] | 2004/5 | Papua New Guinea District Mimika | Moderate stable | All patients attending health facility past 3 days | Routine HMIS surveillance | 1 hospital: n = 186040; 14 primary health clinics: n = 253987 |
| Mathanga et al. [42] | 2005 | Malawi District Phalombe | Moderate stable | Children living in households, aged 6–30 months | Systematic | 12 chronic care clinics: n = 1167 |
| Mathanga et al. [42] | 2010 | Malawi District Phalombe | Moderate stable | All children attending health facility in the past 3 days | Systematic | 12 chronic care clinics: n = 1167 |

(Continued)
| Study                          | Study year(s) | Country   | Geographic unit of comparison | Site(s)                  | Malaria endemicity* | EAG Participants | Sampling methods | Sampling Units; No. sampled | Population Participants | Sampling methods | Sampling Units; No. sampled |
|-------------------------------|---------------|-----------|-------------------------------|--------------------------|---------------------|-------------------|------------------|-----------------------------|-------------------------|-------------------|----------------------------|
| Ndyomugyenyi et al. 2007 [47] | 2005          | Uganda    | District Hoima                | Intense stable           | Primary school children ≥ 10 years | Purposeful 39 primary schools; n = 3602 | Household heads or spouses | Stratified random | 39 villages; n = 2798 households |
| Oduro et al. 2011 [41]        | 2008          | Gambia    | Country Ableda Kaur Yorobawol Gambisara Bureng Gunjur | Moderate stable-         | All patients attending health facilities | Successive 6 health centres; n1 = 4543 (rainy/post rainy season) n2 = 4101 (dry season) | All villagers | Age-stratified random | 18 villages (3 from each catchment area); n1 = 3870 households (rainy/post rainy season) n2 = 3716 households (dry season) |
| Skarbinski et al. 2008 [40]  | 2005          | Tanzania  | Region Lindi                 | Intense stable-          | Children < 5 years coming for sick and well child visits | Stratified cluster sampling (Lindi) | Household members | Stratified random, probability proportional to enumeration area (Lindi) | 22 enumeration areas; n = 574 households |
|                              |               |           | District Rufiji              |                          |                     |                   |                  | Simple random (Rufiji) | N/A; n = 673 households |
| Stevenson et al. 2013 [48]   | 2010          | Kenya     | District Rachuonyo Kisii     | Moderate stable-         | Primary school children in classes 2–6 | 46 government primary schools; n = 4888 | Gender-stratified random sampling | All children > 6 months living in compounds | Simple random sampling, within 600m of each school | N/A; n = 3472 households |

*Malaria endemicity:
Moderately stable endemicity: PfPR = 5.1–39.99% i.e. hypo-mesoendemic
Intensely stable endemicity: PfPR>40% i.e. hyper-holoendemic
Unstable endemic: PfPR <5%.

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Table 4. Comparison of estimates of coverage of control interventions between EAGs and the population.

| Control intervention coverage | Type of EAG survey | EAG survey | Population survey | Fisher’s exact p-value |
|-------------------------------|--------------------|------------|------------------|------------------------|
|                               | Events (n/N)       | Percentage prevalence (95% CI) | Events (n/N)       | Percentage prevalence (95% CI) |
| Household bed net ownership   |                    |            |                  |                        |
| Briand et al                  |                    |            |                  |                        |
| • Salavan, Laos               | ANC                | 307/331    | 92.8 (90.0; 95.5) | 204/205 99.5 (98.5; 100.0) | <0.001 |
| Ndomoyugeni et al             |                    |            |                  |                        |
| • Hoima, Uganda               | School             | 1261/3602  | 35.0 (33.5; 36.6) | 867/2798 30.9 (29.3; 32.7) | <0.001 |
| SkarbINKski et al             |                    |            |                  |                        |
| • Lindi, Tanzania             | Health Facilities  | 506/637    | 79.4 (76.3; 82.6) | 163/354 46.1 (40.9; 51.2) | <0.001 |
| • Rufiji, Tanzania            | Health Centre      | 1195/1433  | 83.4 (81.5; 85.3) | 337/455 74.1 (70.0; 78.1) | <0.001 |
| Household ITN ownership       |                    |            |                  |                        |
| Ndomoyugeni et al             |                    |            |                  |                        |
| • Hoima, Uganda               | School             | 814/3602   | 22.5 (21.2; 24.0) | 629/2798 22.5 (20.9; 24.0) | 0.9759 |
| Individual bed net use        |                    |            |                  |                        |
| Briand et al                  |                    |            |                  |                        |
| • Salavan, Laos               | ANC                | 305/331    | 92.2 (89.3; 95.0) | 204/205 99.5 (98.5; 100.0) | <0.001 |
| GahUTU et al                  |                    |            |                  |                        |
| • Butare, Rwanda              | Health Centre      | 71/102     | 69.6 (60.7; 78.5) | 286/543 52.7 (48.5; 56.9) | 0.0016 |
| • Butare, Rwanda              | Hospital           | 74/102     | 72.6 (63.9; 81.2) | 286/543 52.7 (48.5; 56.9) | <0.001 |
| Mathanga et al                |                    |            |                  |                        |
| • Malawi                     | Health Centre      | 671/1637   | 41.0 (38.6; 43.4) | 420/926 45.4 (42.2; 48.6) | 0.0339 |
| • Malawi                     | Health Centre      | 1067/1909  | 55.9 (53.7; 58.1) | 1899/4565 41.6 (40.2; 43.0) | <0.001 |
| Oduro et al                   |                    |            |                  |                        |
| • Gambia (2005)               | Health Centre      | 3568/4543  | 78.5 (77.3; 79.7) | 3348/3870 86.5 (85.4; 87.6) | <0.001 |
| • Gambia (2008)               | Health Centre      | 2848/4101  | 69.5 (68.0; 70.9) | 2934/3716 79.0 (77.7; 80.3) | <0.001 |
| SkarbINKski et al             |                    |            |                  |                        |
| • Lindi, Tanzania             | Health Facilities  | 507/637    | 79.6 (76.5; 82.7) | 163/354 46.1 (40.9; 51.2) | <0.001 |
| • Rufiji, Tanzania            | Health Centre      | 1195/1463  | 81.7 (79.7; 83.7) | 337/455 74.1 (70.0; 78.1) | <0.001 |
| Stevenson et al               |                    |            |                  |                        |
| • Western Kenya               | School             | 595/1780   | 33.4 (31.2; 35.6) | 2137/3742 57.1 (55.5; 58.7) | <0.001 |

(Continued)
attending health facilities for sick visits were not only concordant with population estimates but there was also concordance between results derived by microscopy and PCR. The accuracy of estimates \( P_f PR \) by RDT (Paracheck \(^R\), Orchid Biomedical Systems, India) derived from school children with circumferential distance was assessed in one study [48], and the estimate from this EAG remained consistently higher with increasing distance within the school catchment area.

Three studies assessed the ability of EAGs to measure changes in \( P_f PR \) as result of changes in coverage of interventions [42, 45] or seasonal transmission [41]. When data was collected before and one year after an ITN campaign in Papua New Guinea, the derived reduction in \( P_f PR \) by RDT in patients with a history of fever attending health facilities (absolute Risk Difference or RD = 23.3%, 95%CI 20.1%, 26.5%) was almost thrice that in the population (RD = 7.6%, 95%CI 6.1%, 9.1%) [45]. After a period of intense distribution of ITNs and a change in first line therapy of malaria from sulphadoxine-pyrimethamine to artemether-lumefantrine in Malawi, the reduction in \( P_f PR \) by malaria microscopy measured in children 6–30 months attending well child clinics (RD = 17.4%, 95%CI 14.6%, 20.2%) was higher than that in the same age strata in the population (RD = 10.5%, 95% CI 7.4%, 13.7%) [41], probably due to significantly higher EAG estimates in the first survey (Table 5). The study by Oduro et al [41] assessed the effect of seasonality on summary estimates \( P_f PR \) by malaria microscopy in all patients attending HFs in six ecologically diverse areas in Gambia, a country with intensely seasonal malaria transmission. In patients attending health facility regardless of cause, the reduction in \( P_f PR \) between the rainy season and the dry season (RD = 22.8%, 95%CI 21.6%, 24.1%) was almost twice that from the HF catchment population (RD = 10.4%, 95%CI 9.3%, 11.6%), probably due to the significantly higher estimates in the rainy/post-rainy season.

Two studies compared estimates of anaemia prevalence rate (APR) between EAGs and the population [41, 42]. In the study by Mathanga et al [42], estimates of APR from children attending well child clinics were not only concordant with values derived from the same age strata in the population but this metric in children attending well child clinics (RD = 2.8%, 95% CI 0.4%, 5.3%) accurately reflected the reduction in the population (RD = 5.3%, 95% CI 2.6%, 8.1%). The other study in Gambia assessed the impact of seasonality on estimates of APR derived from patients of all ages [41], and the difference between the rainy and dry season

### Table 4. (Continued)

| Control intervention coverage | Type of EAG survey | EAG survey | Population survey | Fisher's exact p-value |
|-----------------------------|------------------|------------|------------------|------------------------|
| Mathanga et al              | Health Centre   |            |                  |                        |
| • Malawi\(^d\)              | 601/1637        | 36.7 (34.4; 39.1) | 380/926 | 41.0 (37.9; 44.2) | 0.0311               |
| • Malawi\(^d\)              | 943/1909        | 49.4 (47.2; 51.6) | 1703/4565 | 37.3 (35.9; 38.7) | <0.001               |
| Skarbinski et al            | Health Facilities |            |                  |                        |
| • Lindi, Tanzania           | 245/637         | 38.5 (34.7; 42.2) | 78/354 | 22.0 (17.7; 26.4) | <0.001               |
| • Rufiji, Tanzania          | 1042/1433       | 72.0 (70.4; 75.0) | 241/455 | 53.0 (48.4; 57.6) | <0.001               |
| IRS coverage                |                  |            |                  |                        |
| Stevenson et al             | School          |            |                  |                        |
| • Western Kenya             | 1216/1780       | 68.3 (66.2; 70.5) | 2762/3742 | 73.8 (72.4; 75.2) | <0.001               |

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### Table 5. Comparison of estimates of coverage of malaria morbidity between EAGs and the population.

| Control intervention coverage | Type of EAG survey | EAG survey | Population survey | Fisher’s exact p-value |
|-------------------------------|-------------------|------------|-------------------|------------------------|
| PPR                           |                   |            |                   |                        |
| Gahutu et al                  |                   |            |                   |                        |
| • Butare, Rwanda (BS) Health Centre | 17/103          | 16.5 (9.3; 23.7) | 61/545          | 11.2 (8.6; 13.8)       | 0.1286                |
| • Butare, Rwanda (BS) Hospital | 10/101           | 9.9 (4.1; 15.7)  | 61/545          | 11.2 (8.6; 13.8)       | 0.8625                |
| • Butare, Rwanda (PCR) Health Centre | 22/103          | 21.4 (13.4; 29.3) | 88/545          | 16.2 (13.1; 19.2)      | 0.1994                |
| • Butare, Rwanda (PCR) Hospital | 15/101           | 14.9 (7.9; 21.8) | 88/545          | 16.2 (13.1; 19.2)      | 0.8824                |
| Hetzel et al                  |                   |            |                   |                        |
| • Momase and Highlands, Papua New Guinea (RDT) Health Centre | 402/1304         | 30.8 (28.3; 33.3) | 199/1967        | 10.1 (8.8; 11.5)       | <0.001                |
| • Momase and Highlands, Papua New Guinea (RDT) Health Centre | 50/667           | 7.5 (5.5; 9.5)   | 50/1986          | 2.5 (1.8; 3.2)         | 0.001                 |
| Karyana et al                 |                   |            |                   |                        |
| • Mimika, Papua New Guinea (BS) Health Centre | 36848/253987    | 14.5 (14.4; 14.7) | 290/3890        | 7.5 (6.6; 8.3)         | <0.001                |
| • Mimika, Papua New Guinea (BS) Hospital | 16895/168217    | 10.0 (9.9; 10.2)  | 290/3890        | 7.5 (6.6; 8.3)         | <0.001                |
| • Mimika, Papua New Guinea (BS) Hospital | 4195/17823      | 23.5 (22.9; 24.2) | 290/3890        | 7.5 (6.6; 8.3)         | <0.001                |
| Mathanga et al                |                   |            |                   |                        |
| • Malawi (2005, BS) Health Centre | 464/1516        | 30.6 (28.3; 32.9) | 195/799         | 24.4 (21.3; 27.4)      | 0.0017                |
| • Malawi (2008, BS) Health Centre | 247/1871        | 13.2 (11.7; 14.7) | 607/4377        | 13.9 (12.8; 15.0)      | 0.4945                |
| Oduro et al                   |                   |            |                   |                        |
| • Gambia (BS) Health Centre | 1088/4543       | 24.0 (22.7; 25.2) | 487/3870        | 12.4 (11.3; 13.4)      | <0.001                |
| • Gambia (BS) Health Centre | 46/4101         | 1.1 (0.8; 1.4)   | 80/3716         | 2.2 (1.7; 2.6)         | <0.001                |
| Stevenson et al               |                   |            |                   |                        |
| • Western Kenya School | 454/1780        | 25.5 (23.5; 27.5) | 580/3742        | 15.5 (14.3; 16.7)      | <0.001                |
| APR                           |                   |            |                   |                        |
| Mathanga et al                |                   |            |                   |                        |
| • Malawi (2005) Health Centre | 299/1636        | 18.3 (16.4; 20.2) | 184/926         | 19.9 (17.3; 22.4)      | 0.3440                |
| • Malawi (2008) Health Centre | 295/1909        | 15.5 (13.8; 17.1) | 649/4461        | 14.6 (13.5; 15.6)      | 0.3557                |
| Oduro et al                   |                   |            |                   |                        |
| • Gambia Health Centre | 440/4400        | 10.0 (9.1; 10.9)  | 283/3824        | 7.4 (6.6; 8.2)         | <0.001                |
| • Gambia Health Centre | 317/3963        | 8.0 (7.2; 8.8)   | 127/3716        | 3.4 (2.8; 4.0)         | <0.001                |
| AbPR                          |                   |            |                   |                        |

(Continued)
estimates from this EAG (RD = 2.0%, 95%CI 0.8%, 3.2%) was similar to that in the population (RD = 4.0%, 95%CI 3.0%, 5.0%) though both EAG estimates were consistently higher than population estimates (Table 5).

Two studies compared estimates of antibody prevalence between EAGs and the population [41, 48]. In the study in Gambia where malaria is intensely seasonal with one seasonal peak [41], the difference in Merozoite Surface Protein 19 (MSP1<sub>19</sub>) seroprevalence between the seasons in the EAG (RD = 12.5%, 95%CI 10.4%, 14.6%) was higher than the population (RD = -0.1%, 95%CI -2.0%, 1.8%) due to overestimation of the population value in the rainy season (Table 5). In a moderately stable malaria transmission setting where there are two seasonal peaks of transmission (one major and the other minor), an assessment of AbPR using a number of antigens including Apical Membrane Antigen 1 (AMA1) and MSP1<sub>48</sub> in the month immediately after the major peak revealed that the estimate from school children (AbPR = 51.5%, 95% CI 49.2%, 53.8%) was concordant with that of the same age strata in the population (AbPR = 51.5%, 95% CI 49.9%, 53.1%, p = 1.000), and remained so with increasing distance within the school catchment area.

**Assessment of accuracy of EAG estimates**

Except for the study by Ndyomugyenyi et al [47] were estimates of household ITN ownership derived from primary school children accurately represented population coverage (RD = 0, 95% CI -0.02, 0.02, p = 0.9759), estimates of control intervention coverage derived from EAGs were subject to widely varying degrees of inaccuracy (RD range: -0.24–0.42), with EAGs estimates more commonly but not exclusively over-estimating population values (Fig 2). In the two studies that assessed the accuracy of multiple indicators of intervention coverage [40, 42], estimates of related indicators usually had a similar level of inaccuracy (Fig 2). In the study by Mathanga et al [42], serial estimates of control intervention exhibited similar degree of bias in estimates of individual bed net and ITN use in the first survey but were markedly different in the subsequent survey (Fig 2). In the study by Skarbinski et al [40], the degree of accuracy in estimates of household bed net ownership, individual bed net and ITN use was the same for both well and sick child visits in Rufiji and ITN use in Lindi, whilst estimates of household bed net ownership and individual bed net use were much higher in Lindi (Table 2, Fig 2) indicating regional-specific bias (Fig 2).

### Table 5. (Continued)

| Control intervention coverage | Type of EAG survey | EAG survey | Population survey | Fisher’s exact p-value |
|------------------------------|-------------------|------------|-------------------|-----------------------|
|                              | Events (n/N)      | Percentage prevalence (95% CI) | Events (n/N) | Percentage prevalence (95% CI) |
| Oduro et al                  |                   |            |                   |                       |
| • Gambia                     | Health Centre     | 1122/3380  | 33.2 (31.6; 34.8) | 736/3522              | 20.9 (19.6; 22.2)     | <0.001               |
| • Gambia                     | Health Centre     | 696/3362   | 20.7 (19.3; 22.1) | 712/3391              | 21.0 (19.6; 22.4)     | 0.7875               |
| Stevenson et al             |                   |            |                   |                       |
| • Western Kenya              | School            | 2536/4888  | 51.5 (49.2; 53.8) | 1927/3742             | 51.5 (49.9; 53.1)     | 1.0000               |

BS = Blood slide
PCR = Polymerase chain reaction
RDT = Rapid diagnostic test.

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Estimates of PfPR were on average more consistent than estimates of intervention coverage (Fig 2). In the study by Gahutu et al [44], estimates of PfPR by microscopy and PCR derived from EAGs at different health facility levels were concordant with population values (Fig 2). In the study by Mathanga et al [42], though serial estimates of PfPR from children aged 6–30 months attending well child clinics accurately detected transmission reduction in the same age strata in the population, the estimate of PfPR from this EAG was slightly higher than that in the population in 2005 (RD = 0.06, 95% CI 0.02, 0.10, p = 0.002). Estimates of APR derived from EAGs in two studies [41, 42] were overall a more consistent estimation of population prevalence than PfPR (Fig 2). The close approximation of EAG estimates of APR together with its accurate measurement of a reduction in population prevalence suggests that it is a good surrogate indicator for APR in the population [42]. Estimates of AbPR derived from EAGs were more accurate in the dry season in the Gambia [41], with rainy season estimates being higher than population estimates (RD = 0.12, 95% CI 0.10, 0.02, p<0.001).

Two of the studies attempted to measure the degree of inaccuracy or improve the precision of estimates by controlling for bias [40, 42]. After controlling for potential confounders (age in months, child’s sex, survey type and study area) in a multivariable analysis, in the study by Skarbinski et al [40], the adjusted odds ratio (aOR) between the health facility survey and the EAG survey for individual bed net use (aOR = 2.05, 95% CI 1.36, 3.08) and ITN use (aOR = 2.41, 95% CI 1.69, 3.44) still indicated an overestimation of population coverage. In the study by Mathanga et al [42], after adjusting for confounders in a multivariate analysis, parasitaemia in 2008 vs 2005 in children attending well child clinic (aOR = 0.31, 95% CI 0.22, 0.46) was equivalent to that in the same age strata in the population (aOR = 0.40, 95% CI 0.30, 0.52), and this was similar for anaemia (Hb<8.0d/dl) in this EAG (aOR = 0.85, 95% CI 0.65, 1.65) compared to the population (aOR = 0.74, 95% CI 0.59, 0.94).

For EAG to guide control efforts, it should correctly classify the uptake of control interventions and malaria endemicity. The prevalence difference in bed net use suggested that EAG surveys overestimated population levels up to a certain point (population coverage of approximately 72%), after which they overestimated population values, but this trend was not statistically significant (p = 0.993) (Fig 3A). The prevalence difference in PfPR overestimated population prevalence with increasing transmission (p = 0.979) (Fig 3B), but our assumptions are also limited by the fact that the studies included in this review only covered moderately stable and unstable endemic transmission intensities. Based on the classification of malaria endemicity from the PfPR results, most of the EAG surveys (13/14) were concordant with that of the population (Table 6). During the post-ITN survey in Papua New Guinea [45], population PfPR dropped to unstable endemic levels (PfPR = 2.5%, 95%CI 1.8%; 3.2%) but was wrongly classified to be moderate stable by the EAG (PfPR = 7.5%, 95%CI 5.5%; 9.5%).

Discussion

Monitoring control progress is important to assess the effectiveness and coverage of malaria control programmes. Easy access group surveys are easier to conduct than population surveys and could provide accurate monitoring of control progress if the EAG sample is representative of our population stratum of interest [30, 33, 50–53]. Review of the available literature on EAGs suitable for district or sub-district surveillance of malaria control progress revealed a wide variation in the precision of estimates between and within studies, particularly for estimates of control intervention coverage. The small number of studies in this review shows how little effort has been made to explore the potential approach and settings for use of EAGs, probably due to the inherent assumption of bias in such opportunistic samples. Our study has potential limitations. Our search strategy may not have identified all the relevant papers or
### Surveillance in EAGs as a tool for evaluating malaria control progress

**Study and year** | **RD (95% CI)**<br>
--- | ---
**Household bednet ownership**<br>Briand 2014 (ANC) | -0.07 (-0.10, -0.04)<br>Ndyomugenyi 2005 (SC) | 0.04 (0.02, 0.06)<br>Skarbinski 2005 (WSC) | 0.09 (0.05, 0.14)<br>Skarbinski 2005 (WCC) | 0.33 (0.27, 0.39)

**Household ITN ownership**<br>Ndyomugenyi 2005 (SC) | 0.00 (-0.02, 0.02)

**Individual bednet use**<br>Briand 2014 (ANC) | -0.07 (-0.10, -0.04)<br>Gahutu 2010 (WSC) | 0.17 (0.07, 0.27)<br>Gahutu 2010 (WCC) | 0.20 (0.10, 0.30)<br>Mathanga 2005 (WSC) | -0.04 (-0.08, -0.00)<br>Mathanga 2008 (WSC) | 0.14 (0.12, 0.17)<br>Odoro 2009 (OPD) | -0.08 (-0.10, -0.06)<br>Odoro 2009 (OPD) | -0.10 (-0.11, -0.08)<br>Skarbinski 2005 (WSC) | 0.34 (0.27, 0.40)<br>Skarbinski 2005 (WCC) | 0.08 (0.03, 0.12)<br>Stevenson 2010 (SC) | -0.24 (-0.26, -0.21)

**Individual ITN use**<br>Mathanga 2005 (WSC) | -0.04 (-0.08, -0.00)<br>Mathanga 2008 (WSC) | 0.12 (0.09, 0.15)<br>Skarbinski 2005 (WSC) | 0.20 (0.15, 0.25)<br>Skarbinski 2005 (WCC) | 0.16 (0.11, 0.22)

**IRS coverage**<br>Stevenson 2010 (SC) | -0.05 (-0.08, -0.03)

**P1PR**<br>Gahutu 2010 (WSC) BS | -0.01 (-0.08, 0.05)<br>Gahutu 2010 (WSC) BS | 0.05 (-0.02, 0.13)<br>Gahutu 2010 (WSC) PCR | 0.05 (-0.03, 0.14)<br>Gahutu 2010 (WSC) PCR | -0.01 (-0.09, 0.06)<br>Hetzl 2008/9 (OPD) RDT | 0.21 (0.18, 0.24)<br>Hetzl 2009/10 (OPD) RDT | 0.05 (0.03, 0.07)<br>Karyana 2004/5 (OPD) BS | 0.03 (0.02, 0.03)<br>Karyana 2004/5 (OPD) BS | 0.07 (0.06, 0.08)<br>Karyana 2004/5 (OPD) BS | 0.16 (0.15, 0.17)<br>Mathanga 2005 (WSC) BS | 0.06 (0.02, 0.10)<br>Mathanga 2008 (WSC) BS | -0.01 (-0.03, 0.01)<br>Odoro 2009 (OPD) BS | -0.01 (-0.02, -0.00)<br>Odoro 2009 (OPD) BS | 0.12 (0.10, 0.13)<br>Stevenson 2010 (SC) RDT | 0.10 (0.08, 0.12)

**APR**<br>Mathanga 2005 (WSC) | -0.02 (-0.05, 0.02)<br>Mathanga 2008 (WSC) | 0.01 (-0.01, 0.03)<br>Odoro 2009 (OPD) | 0.04 (0.03, 0.05)<br>Odoro 2009 (OPD) | 0.02 (0.01, 0.04)

**AbPR**<br>Odoro 2009 (OPD) | 0.06 (0.04, 0.07)<br>Odoro 2009 (OPD) | -0.02 (-0.04, -0.00)<br>Stevenson 2010 (SC) | 0.00 (-0.03, 0.03)
there may be other sources of grey literature that may have been missed. We phrased our search terms as simply as possible to allow a wider inclusion of possible papers and in this regard, we may have missed some papers with highly selective titles. The studies selected for the systematic review only included health facility (including ANC) and school surveys, and were from settings with moderate and intense stable malaria transmission, so our results may not be applicable to other EAGs or transmission settings. Our literature search was guided by categories of EAGs with historical evidence of use for malaria surveillance or which we theorized would be suitable for malaria surveillance at the district or sub-district level. This may have excluded publications on other potential EAGs. We limited our review to studies that compared EAG samples to populations samples of the same age or other at-risk stratum. Whilst this may improve the accuracy of EAG estimates of PfPR, especially in moderate to severe transmission settings, this does not mean that EAGs could not be used to estimate control intervention coverage in any population stratum or PfPR at the lower end of the transmission spectrum in other population at-risk strata. Given the pace of developments in analytical technics, this is an area where substantial gains can be made and we discuss this below.

Dealing with bias in EAG surveillance

The main cause of bias in EAG surveillance is due to the selection of an unrepresentative sample of the population of interest. The opportunistic nature of the sampling frame in EAGs is inherently susceptible to selection bias when EAG sampling captures an unrepresentative subset of the population of interest. Particularly, if the reason for inclusion in the EAG sample is associated with the indicator of interest. For example, given the fact that those who are wealthier and more educated are more likely to attend health facilities, and have access to or use ITNs [54, 55], self-reported ITN possession and use from health facility surveys is likely to over-estimate ITN coverage in the population. This could be corrected using the verification rate measured from a small random sample of the catchment population. Also, the representativeness of estimates of PfPR from health facilities is likely to be affected by the difference in transmission between malaria seasons, overall malaria transmission and the prevalence of non-malaria fevers. This could be limited by the use of EAGs excluding individuals coming for sick visits [56, 57] or prioritizing indicators that are less sensitive to short-term changes in transmission like AbPR [58, 59]. Population APR is also less sensitive to short term changes in transmission [60], but whether this makes it an appropriate indicator to measure changes in transmission is debatable. Though malaria is an important correlate of anaemia in children, the aetiology of anaemia is multifactorial and in particular the role of other infections, poor nutrition and the interaction between malaria and nutrition needs to be clarified [61]. Where there is a high probability of inclusion in the EAG sample, the difference in the estimates of an indicator measured from individuals who are and are not included in the EAG sample is likely to be less significant, and the EAG sample is more likely to be representative of the true situation in the population. For example, coverage rates of public health interventions were similar between vaccinated and unvaccinated children if population vaccine coverage was over 60% [62]. Most of the standard methods for analysis of data from convenience samples are based on the questionable assumption that selection bias can be exclusively ascribed to measured risk factors for malaria. Novel geostatistical methods have been recently developed to relax this assumption [63]. By combining data from unbiased gold-standard surveys and opportunistic
A. Bednet use

Pearson correlation coefficient = -0.0074, p = 0.993

B. PfPR

Pearson correlation coefficient = -0.0076, p = 0.979
samples, these methods are able to correct for the selection bias in the convenience samples that is induced by both measured and unmeasured risk factors. Though the etiology of health facility access and utilization is multifactorial [64–69], health facility utilization follows a geographic pattern [70–73] and if this can be accurately measured through a small geospatially random sample of the population and accounted for in the model, will allow correction for bias and the production of accurate maps of control progress. Where point estimates are required, combining the EAG sample with a small and presumably far less expensive random sample of the population [74], the so-called hybrid sampling methodology will generate more accurate hybrid prevalence estimates. Pooling data from multiple EAGs in our area of interest is also likely to improve the precision of point estimates [47, 75].

Another cause of bias in EAG samples mainly affecting reported coverage of control interventions is social desirability bias. Survey respondents may answer questions in a manner they consider favourable to the interviewee leading to erroneously high self-reporting of coverage of control interventions [76]. This may be further compounded by the inability to directly validate the presence and use of household-level and individual control measures as in population surveys. Few studies have assessed the effect of social desirability bias on the effect of bed net use [77–80], and the wide range in verification rate of bed net use after self-report (60.9–96.2%) suggests variability in the effect of social desirability bias from setting to setting. Social desirability bias can be limited by modifying the standard MIS questionnaire [13] to include questions aimed at detecting and measuring social desirability bias so it can be directly accounted for in analysis [81], improving interviewee confidence by indicating the anonymity of their responses [81], or indirectly by correcting for the verification rate measured from a random sample of the catchment population.

Use of EAG surveys to measure progress in malaria control

Before EAGs can be routinely used to measure malaria control progress, there are a few issues to address. Firstly, how much inaccuracy we are willing to tolerate? If the purpose of the survey is to measure trends in point estimates of control progress, some degree of inaccuracy is tolerable if EAG data displays similar trends to population data; as evidenced by the successful demonstration of transmission reduction from health facility surveys in some endemic countries [30, 33, 50–53], and increasing endorsement by WHO as a surveillance tool in different transmission settings and phases of control [1]. One study suggested that estimates of population PPR from health facility surveys might misclassify malaria endemicity at the lower end of the transmission spectrum [45], but the population in this study (i.e. all health facility attendees) may not be the most suitable to capture the most at-risk population at low transmission settings. When more accurate point estimates are required or accurate data is required over a large geographic area, pooling data from multiple similar EAGs [47, 75] or hybrid sampling methodology [74] may improve precision. If the purpose of the survey is to measure changes

| Table 6. Relationship between the results of the classification of malaria endemicity between EAG and population sampling. |
|-----------------------------------------------|
| EAG               | Moderate stable | Unstable endemic | Total |
| Moderate stable   | 12              | 1                | 13    |
| Unstable endemic  | 0               | 1                | 1     |
| Total             | 12              | 2                | 14    |

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in the geospatial distribution of uptake of control interventions and transmission intensity, to identify areas of low intervention coverage and potential hotspots respectively for targeted control intervention delivery; the smaller sized EAG catchment areas compared to community-based surveys [70–73] means the maps derived using EAG sampling will not be consistent with those derived using community-level data and would require geospatial statistical methods to correct for bias [63].

Secondly, are EAGs surveys more cost effective than standard approaches? Because of the ease of EAG sampling, conducting an EAG survey should theoretically be cheaper than a population survey in the same catchment area. Reports from school surveys in Kenya seem to suggest that the financial cost of school surveys is less than half that of a household survey [9, 11]. Though a detailed economic costs analysis of school surveys has not been done in comparison to those from household surveys, and the lower financial costs has not been validated in other EAGs; the decreased expenditure on personnel, transportation and communication in school surveys compared to household surveys suggest that surveillance in EAGs is likely to be more cost-effective [9].

Thirdly, when are EAG surveys most likely to be beneficial i.e. to complement malaria programmatic efforts? At moderate to high malaria transmission intensity, surveillance systems rely on passive surveillance (e.g. HMISs) supplemented by large serial populations surveys (e.g. MISs), with data reported at the national, regional and sometimes district level. Surveillance in EAGs in such settings will be beneficial in providing more detailed sub-district estimates from “problematic” districts with poor control progress compared to national average, estimates from hard-to-reach communities (e.g. opportunistic surveys during MDA) who would otherwise not be covered by population surveys, or when data is required to assess at-risk stratum specific control interventions (e.g. ANC and delivery surveys to assess the impact of Intermittent Preventive Treatment in pregnancy or IPTp). These EAG surveys should be carried out at the same time as population surveys i.e. every 2–3 years, so that the estimates can be interpreted within the context of a wider perspective of population control progress. As transmission intensity falls and we approach the elimination phase, reorientation of programmatic efforts are required to identify hotspots [8] and special high-risk populations [82, 83] both of which serve as reservoirs of infection that should be targeted for malaria elimination. Population surveys become less logistically attractive and less practical given the fact that more regular (e.g. quarterly) local (sub-district) level data is required on control progress. Surveillance in EAGs becomes more attractive as a more sustainable method of surveillance including the high-risk groups (e.g. rural community market surveys at border crossings).

Finally, how do we integrate surveillance in EAGs with current control strategies? EAG surveillance can provide timely data of reasonable accuracy on control progress that reflect local variation at the district or sub-district level, and is complementary to national community-based surveys like MISs [13]. EAG surveys can provide a means of rapid assessment of areas known to have poor coverage or key population risk-strata. The ease of sampling and low costs allows more frequent or even continuous surveys providing timely data and encouraging reactive targeted control. EAG surveillance in health facilities may have a motivational impact on health workers at the district and sub-district level through the provision of continuous locally appropriate data on intervention coverage and malaria transmission, and its flexibility allows it to adapt to new programmatic needs over time. Sufficient person-time is however needed for successful data acquisition in health facilities and to ensure no duplication with recurrent data collection. Implementing and scaling up EAG surveillance will require minimal reorientation and structuring of the health system, including determining which health facility personnel should be dedicated to malaria surveillance, and some preparation and buy-in is required by both national and global health players.
Conclusions

This review describes the previous experiences with the validation of estimates of malaria control progress from different EAGs and highlights the potential of surveillance in EAGs as a complementary approach to current surveillance systems. The utility of an EAG for routine surveillance of progress in malaria control at the district or sub-district programmatic level will be driven by several factors including whether serial point estimates or more precise geospatial distribution is required, the degree of precision accepted, the desired population of interest (e.g. at-risk groups), and the resources available for surveillance (both financial and otherwise). The low cost of EAG surveillance, its flexibility and potential to offer locally applicable timely estimates of control which could improve programmatic responses suggest that further validation and optimization is required.

Supporting information

S1 Document. PRISMA 2009 checklist. (DOC)

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Author Contributions

Conceptualization: Sanie S. S. Sesay, Dianne J. Terlouw.
Data curation: Sanie S. S. Sesay.
Formal analysis: Sanie S. S. Sesay.
Funding acquisition: Sanie S. S. Sesay, Dianne J. Terlouw.
Investigation: Sanie S. S. Sesay.
Methodology: Sanie S. S. Sesay, Peter J. Diggle, David Schellenberg, David G. Lalloo, Dianne J. Terlouw.
Project administration: Sanie S. S. Sesay.
Resources: Sanie S. S. Sesay.
Software: Sanie S. S. Sesay.
Supervision: Peter J. Diggle, David G. Lalloo, Dianne J. Terlouw.
Validation: Sanie S. S. Sesay, David G. Lalloo, Dianne J. Terlouw.
Visualization: Sanie S. S. Sesay.
Writing – original draft: Sanie S. S. Sesay, Emanuele Giorgi, David G. Lalloo, Dianne J. Terlouw.
Writing – review & editing: Sanie S. S. Sesay, Emanuele Giorgi, Peter J. Diggle, David Schellenberg, David G. Lalloo, Dianne J. Terlouw.
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