METHOD ARTICLE

Optimisation and standardisation of a multiplex immunoassay of diverse *Plasmodium falciparum* antigens to assess changes in malaria transmission using sero-epidemiology [version 2; peer review: 1 approved, 1 approved with reservations]

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Abstract

**Background:** Antibody responses have been used to characterise transmission and exposure history in malaria-endemic settings for over a decade. Such studies have typically been conducted on well-standardised enzyme-linked immunosorbent assays (ELISAs). However, recently developed quantitative suspension array technologies (qSAT) are now capable of high-throughput and multiplexed screening of up to hundreds of analytes at a time. This study presents a customised protocol for the Luminex MAGPIX® qSAT using a diverse set of malaria antigens. The aim is to develop a standardised assay for routine serological surveillance that is implementable across laboratories and epidemiological settings.

**Methods:** A panel of eight *Plasmodium falciparum* recombinant antigens, associated with long- and short-lived antibody responses, was designed for the Luminex MAGPIX® platform. The assay was optimised for key steps in the protocol: antigen-bead coupling concentration, buffer composition, serum sample dilution, and bead storage conditions. Quality control procedures and data normalisation methods were developed to address high-throughput assay processing. Antigen-specific limits of quantification (LOQs) were also estimated using both in-house and WHO reference serum as positive controls.

**Results:** Antigen-specific bead coupling was optimised across five serum dilutions and two positive controls, resulting in concentrations operational...
dilutions and two positive controls, resulting in concentrations operational within stable analytical ranges. Coupled beads were stable after storage at room temperature (22°C) for up to eight weeks. High sensitivity and specificity for distinguishing positive and negative controls at serum sample dilutions of 1:500 (AUC 0.94 95%CI 0.91-0.96) and 1:1000 (AUC 0.96 95%CI 0.94-0.98) were observed. LOQs were also successfully estimated for all analytes but varied by antigen and positive control.

**Conclusions:** This study demonstrates that developing a standardised malaria-specific qSAT protocol for a diverse set of antigens is achievable, though further optimisations may be required. Quality control and data standardisation methods may also be useful for future analysis of large sero-epidemiological surveys.

**Keywords**
Plasmodium, malaria, antibodies, serum, serology, Luminex, MAGPIX

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Introduction

Until recently, the vast majority of malaria antibody studies have sought to understand the acquisition of protective immunity to inform vaccine development\(^1\). However, there is growing interest in identifying new serological markers of malaria exposure for epidemiological surveillance\(^2\). Rational selection of these markers for population-wide sero-profiling could enable the development of improved tools for monitoring changes in malaria transmission. Such tools have the potential to simultaneously characterise both historical and recent patterns in malaria exposure.

Accurately quantifying antibody dynamics in these contexts requires population-representative or frequently-sampled longitudinal datasets\(^3\). Sero-epidemiological studies of this scale have historically used enzyme-linked immunosorbent assays (ELISAs), which are easily standardised, widely available, and ideal for high-throughput analysis of a single antigen (or the combined response to multiple antigens). However, the limited dynamic range and need for relatively high blood volumes make ELISAs less efficient for evaluating multiple analyte-specific responses. Conversely, protein microarrays allow high-throughput analysis of hundreds to thousands of analytes per sample\(^4\), but are still prohibitively expensive and not easily accessible by national malaria control programmes (NMCPs) or laboratories.

Cytometric bead array (CBA) and quantitative suspension array technologies (qSATS), such as Luminex xMAP\(^\circledR\) (Luminex Corp, Austin TX), are now available as affordable mid- to high-throughput multiplexing platforms. These offer several advantages, including the simultaneous quantification of 50-500 proteins in a single well, the use of standard 96- or 384-well plates, and requiring as little as 5µl of plasma or serum\(^5\). These platforms have also been shown to measure a larger dynamic range of antibody responses compared to ELISA\(^6\). Optimisation of CBA and qSAT platforms can be complex for antigen panels designed to capture a wide range of antibody dynamics. A key challenge is achieving a high degree of multiplexing while retaining differential responses across a diverse set of antigens\(^1\). Additionally, standardised approaches for the epidemiological analysis of Luminex data are still in development. A number of recent studies have assessed cluster-level antibody responses based on Luminex data for malaria\(^1\) and other infectious diseases\(^8\), but there is still a paucity of data on appropriate methods for standardisation and interpretation across laboratories, sites, and antigens.

In this study, we developed a customised panel of *Plasmodium falciparum* (*Pf*) recombinant antigens as serological markers of both historical and recent malaria exposure and optimised a protocol for the Luminex MAGPIX\(^\circledR\) qSAT platform. This includes five recently developed antigens previously validated in protein microarray studies for their association with recent malaria infection in Ugandan and Malian children\(^1\). For epidemiological analysis, we present quality control procedures for high-throughput assay processing, data normalisation methods, and report estimates of antigen-specific limits of quantification (LOQs).

The aim was to translate the development of a suite of markers for malaria exposure to a qSAT platform that is practical for epidemiological surveillance across laboratories and countries.

Methods

Assay conditions were assessed and optimised for key steps in the protocol: antigen-bead coupling concentration, buffer composition to reduce non-specific reactivity, serum sample dilution, and the impact of storage length and temperature on bead stability (Figure 1).

Antigen selection and design

A multiplex panel was developed for the Luminex MAGPIX\(^\circledR\) suspension bead array containing eight erythrocytic *Pf* recombinant proteins (Table 1). Antigens were selected from an initial screen of 856 candidates on an *in vitro* transcription and translation (IVTT) protein microarray assay based on their correlation with previous malaria infection in children\(^1\). Each antigen was generated and expressed in *Escherichia coli* as glutathione S-transferase (GST)-tagged fusion proteins using methods as previously described by Herman *et al.*,\(^1\) Tetteh *et al.*,\(^1\) and Polley *et al.*\(^1\). The exception to the panel was *PfAMA1*, which was expressed in *Pichia pastoris* as a histidine-tagged protein\(^1\). Protein purification was conducted by affinity chromatography (Glutathione Sepharose 4B (GE Healthcare Life Sciences) or HisPur Ni-NTA (Invitrogen) resins for GST and His tagged proteins, respectively), and the concentration, quality, and purity of the antigen yield was assessed using a Bradford assay and SDS-PAGE. Bacterial lysate was generated from the culture of untransformed *E. coli* and used in the preparation of assay buffers as non-specific protein to eliminate background reactivity to *E.coli* proteins\(^1\).

Positive and negative controls for assay optimisation

For the optimisation of the assay protocol, several different positive controls were used and are summarised for each test
Figure 1. Scheme describing the qSAT assay protocol. Assay conditions tested for optimisation indicated in green boxes and red text.

Table 1. Summary of antigens in multiplex Luminex panel.

<table>
<thead>
<tr>
<th>Gene ID</th>
<th>Antigen name</th>
<th>Allele</th>
<th>Expression</th>
<th>Location</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>PF3D7_0930300</td>
<td>PfMSP1</td>
<td>19</td>
<td>GST</td>
<td>Merozoite surface</td>
<td>19kDa fragment of MSP1 molecule</td>
</tr>
<tr>
<td>PF3D7_1133400</td>
<td>PfAMA1</td>
<td>FVO</td>
<td>His6</td>
<td>Merozoite / Sporozoite</td>
<td>Apical membrane antigen</td>
</tr>
<tr>
<td>PF3D7_1035300</td>
<td>PfGLURP.R2</td>
<td>F32</td>
<td>N/A</td>
<td>Merozoite</td>
<td>Glutamate rich protein R2</td>
</tr>
<tr>
<td>PF3D7_0532100</td>
<td>Etramp5.Ag1</td>
<td>3D7</td>
<td>GST</td>
<td>iRBC/PVM</td>
<td>Early transcribed membrane protein</td>
</tr>
<tr>
<td>PF3D7_0423700</td>
<td>Etramp4.Ag2</td>
<td>3D7</td>
<td>GST</td>
<td>iRBC/PVM</td>
<td>Early transcribed membrane protein</td>
</tr>
<tr>
<td>PF3D7_0402400</td>
<td>GEXP18</td>
<td>3D7</td>
<td>GST</td>
<td>Gametocytes</td>
<td>Gametocyte exported protein</td>
</tr>
<tr>
<td>PF3D7_0501100.1</td>
<td>HSP40.Ag1</td>
<td>3D7</td>
<td>GST</td>
<td>iRBC/Gametocytes</td>
<td>Heat shock protein 40, type II</td>
</tr>
<tr>
<td>PF3D7_1002000</td>
<td>Hyp2</td>
<td>3D7</td>
<td>GST</td>
<td>iRBC/PVM</td>
<td>Plasmodium exported protein</td>
</tr>
<tr>
<td>--</td>
<td>GST</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>GST expression tag</td>
</tr>
<tr>
<td>--</td>
<td>TT</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>Tetanus Toxoid</td>
</tr>
</tbody>
</table>

*Conformational protein. iRBC, infected red blood cell; PVM, parasitophorous vacuole membrane; GST, glutathione S–transferase.
condition (Table 2). This included pooled serum from hyperimmune individuals in Tanzania (CP3), Uganda (PRISM), The Gambia (Brefet) as well as the WHO reference serum (NIBSC 10/198). Individual plasma samples from European malaria-naïve adults were used as negative controls.

**Microsphere coupling, buffer, and sample dilution optimisation**

Recombinant antigens were coupled to MagPlex® COOH-microspheres or ‘beads’ (Luminex Corp., Austin TX) following the protocol described by the Luminex Corporation. Optimal coating concentrations for each antigen were tested using a six-point serial titration of protein. Starting dilutions were determined by the known immunogenicity range of each antigen (Table 3). Titrations were tested under a series of conditions to assess the variability across 1) two different positive pools (CP3 and WHO NIBSC 10/198) and 2) five serum sample dilutions (1:100, 1:200, 1:400, 1:800, and 1:1600). All samples were incubated overnight in buffer B with *E. coli* lysate. The antigen titration at mid-point of the median fluorescence intensity (MFI) dose-response curve (EC₅₀ or MFIₕₕ) was calculated for each positive pool and serum dilution condition (15 in total for each antigen), using Equation 1 described further below. The median titration across all conditions was selected as the optimal coupling concentration for each antigen and used for large volume bead coupling. Beads were then re-suspended in 1 mL of storage buffer (1x phosphate buffered saline (PBS pH 7.2), 0.05% Tween, 0.5% BSA, 0.02% sodium azide, 0.02% Pefabloc (Sigma)) and stored at 4°C until further use.

Antigen reactivity was tested for sensitivity to buffer composition and serum sample dilution. To reduce non-specific background reactivity for antigens expressed in *E. coli*, factorial testing of the two buffer solutions (buffer A - 1xPBS, 0.05% Tween, 0.5% BSA, 0.02% sodium azide; and buffer B - 1xPBS, 0.05% Tween, 0.5% BSA, 0.02% sodium azide, 0.1% casein, 0.5% polyvinyl alcohol (PVA), 0.5% polyvinyl pyrrolidone (PVP)) was conducted. Both buffers were tested with or without supplementation of *E. coli* lysate (added at 15.25 μg/ml), resulting in a total of four different buffer compositions. Positive control samples were 40 individuals from Brefet, The Gambia and negative controls were 40 malaria-naïve European individuals (all samples tested at 1:100 dilution). The effects of buffers A and B - with or without *E. coli* - on MFI values of both positive and negative samples were assessed using linear regression with an interaction term to test for potentially synergistic effects on background reactivity by adding *E. coli* to buffer B.

A range of serum concentrations was also tested (1:100, 1:500, 1:1000 and 1:2000) using samples from 20 individuals from endemic regions in Uganda, Tanzania, and The Gambia as positive controls and 20 malaria naïve European individuals as negative controls. All samples were incubated overnight in buffer B with *E. coli* lysate. Optimal serum sample dilutions were selected based on cross-validated Area Under the Receiver Operating Characteristics Curve (AUC) values, calculated from the sensitivity and specificity of continuous MFI values for predicting a positive or negative control using the *ci.cvAUC* function in the ‘cvAUC’ package version 1.1.0 (R version 3.5.1).

### Table 2. List of assay conditions tested.

<table>
<thead>
<tr>
<th>Optimisation</th>
<th>Values tested</th>
<th>Samples</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Antigen concentration</strong></td>
<td>6-dilution serial protein titration</td>
<td>• Tanzanian pooled serum (CP3)</td>
</tr>
<tr>
<td></td>
<td>2 positive controls</td>
<td>• WHO reference serum (NIBSC 10/198)</td>
</tr>
<tr>
<td></td>
<td>5 serum sample dilutions (1:100, 1:200, 1:400, 1:800, 1:1600)</td>
<td>• 40 Gambian individuals (Brefet)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• 40 malaria naïve European blood donors (PHE)</td>
</tr>
<tr>
<td><strong>Buffer composition</strong></td>
<td>Buffer A</td>
<td>Samples tested at serum dilution of 1:100</td>
</tr>
<tr>
<td></td>
<td>• With <em>E. coli</em> lysate</td>
<td>• Tanzanian pooled serum (CP3)</td>
</tr>
<tr>
<td></td>
<td>• Without <em>E. coli</em> lysate</td>
<td>• 2 sets of Ugandan pooled serum (Apac, PRISM)</td>
</tr>
<tr>
<td></td>
<td>Buffer B</td>
<td>• Gambian pooled serum (Brefet)</td>
</tr>
<tr>
<td></td>
<td>• With <em>E. coli</em> lysate</td>
<td>• 12 Ugandan individuals</td>
</tr>
<tr>
<td></td>
<td>• Without <em>E. coli</em> lysate</td>
<td>• 4 Gambian individuals</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• 20 malaria naïve European blood donors (PHE)</td>
</tr>
<tr>
<td><strong>Sample dilution</strong></td>
<td>1:100, 1:500, 1:1000, 1:2000</td>
<td>Samples incubated in buffer B with <em>E. coli</em></td>
</tr>
<tr>
<td><strong>Microsphere storage</strong></td>
<td>• Length of time</td>
<td>• Tanzanian pooled serum (CP3)</td>
</tr>
<tr>
<td></td>
<td>&lt;4 weeks storage at 4°C</td>
<td>• 2 sets of Ugandan pooled serum (Apac, PRISM)</td>
</tr>
<tr>
<td></td>
<td>6 months storage at 4°C</td>
<td>• Gambian pooled serum (Brefet)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• 4 malaria naïve European blood donors (PHE)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Samples tested at serum dilution of 1:100 and incubated in buffer B with *E. coli</td>
</tr>
</tbody>
</table>
Table 3. Antigen coupling concentration titrations. Optimal range and final antigen concentration in bold.

<table>
<thead>
<tr>
<th>Antigen</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>Final coupling concentration (µg/mL)</th>
<th>Final coupling concentration (ng/5000 beads)</th>
</tr>
</thead>
<tbody>
<tr>
<td>P<em>MSP1</em>19</td>
<td>800.00</td>
<td>100.00</td>
<td>12.50</td>
<td>1.563</td>
<td>0.195</td>
<td>0.024</td>
<td>51.5</td>
<td>20.6</td>
</tr>
<tr>
<td>PIAMA1</td>
<td>600.00</td>
<td>75.00</td>
<td>9.375</td>
<td>1.172</td>
<td>0.147</td>
<td>0.018</td>
<td>3.85</td>
<td>1.54</td>
</tr>
<tr>
<td>PKLURPR2</td>
<td>300.00</td>
<td>37.50</td>
<td>4.688</td>
<td>0.586</td>
<td>0.073</td>
<td>0.009</td>
<td>0.042</td>
<td>0.017</td>
</tr>
<tr>
<td>Etramp5.Ag1</td>
<td>1000.00</td>
<td>125.00</td>
<td>15.625</td>
<td>1.953</td>
<td>0.244</td>
<td>0.031</td>
<td>243</td>
<td>97.2</td>
</tr>
<tr>
<td>GEXP18</td>
<td>750.00</td>
<td>93.75</td>
<td>11.719</td>
<td>1.465</td>
<td>0.183</td>
<td>0.023</td>
<td>618</td>
<td>247.2</td>
</tr>
<tr>
<td>HSP40.Ag1</td>
<td>800.00</td>
<td>100.00</td>
<td>12.50</td>
<td>1.563</td>
<td>0.195</td>
<td>0.024</td>
<td>91.5</td>
<td>36.6</td>
</tr>
<tr>
<td>Etramp4.Ag2</td>
<td>1000.00</td>
<td>125.00</td>
<td>15.625</td>
<td>1.953</td>
<td>0.244</td>
<td>0.031</td>
<td>32.5</td>
<td>13.0</td>
</tr>
<tr>
<td>Hyp2</td>
<td>350.00</td>
<td>43.75</td>
<td>5.469</td>
<td>0.688</td>
<td>0.085</td>
<td>0.011</td>
<td>197</td>
<td>78.8</td>
</tr>
</tbody>
</table>

Stability and reproducibility testing
To evaluate the impact of storage temperature, accelerated stability testing was conducted using aliquots of antigen-coupled beads stored for 1–8 weeks at -20°C, 22°C (room temperature), 37°C and 42°C. Antigen-coupled beads were incrementally added to each storage temperature at intervals over an 8-week period, such that total storage time ranged from 1, 2, 4, and 8 weeks total. All the antigen-coupled beads were run simultaneously at the end of eight weeks with a single positive control titration, avoiding the need to adjust for random effects due to week or positive control batch in subsequent regression analysis. Beads were assayed using a five-point serial dilution titration and the CP3 Tanzanian positive pool (described above) at a 1:100 dilution in buffer B with *E. coli* lysate. The effect of each storage condition was assessed with multivariate linear regression to estimate change in MFI over time, adjusted for storage temperature and sample concentration and allowing for pairwise interaction between all covariates (storage temperature, storage time and sample concentration).

To assess the impact of long-term storage on stability and reproducibility of results, two sets of antigen-coupled beads were assessed; one batch stored for 6 months at 4°C and another batch stored for less than 4 weeks at 4°C. Pooled serum of hyper-immune individuals from Tanzania, Uganda, and The Gambia were each tested in triplicate with beads from each storage condition (less than 4 weeks and 6 months). Change in logMFI at 6 months compared to less than 4 weeks storage was assessed using linear regression, adjusting for antigen and allowing for random effects by sample and replicate.

Final qSAT assay procedure
General assay procedures were as follows and illustrated in Figure 1. First, an initial mixture containing 8 µl of each set of antigen-coupled microspheres and 5 ml of buffer A was prepared, yielding approximately 1,000 beads per region per well (based on optimal conditions reported in previous studies[1–3]). Next, 50 µl of this combined microsphere mixture was added to a 96-well flat bottom plate (BioPlex Pro™, Bio-Rad Laboratories, UK) and washed once by placing the assay plate onto a magnetic plate separator (Bio-Plex®, Bio-Rad Laboratories, UK) and pausing for 2 minutes. Plates were then inverted forcefully to remove the liquid and 100 µl of PBS-T (1xPBS, 0.05% Tween-20) added to each well. Next, 50 µl of samples and controls were added to the plate and incubated in the dark at room temperature (RT) on a microplate shaker at 500 rpm for 90 minutes. Following three washes, 50 µl of fluorescent secondary antibody (Jackson Immuno 109-116-098; Goat anti-human Fcy-fragment specific IgG conjugated to R-Phycocerythrin (R-PE)), diluted to a 1:200 dilution with buffer A, was added to all wells and incubated for 90 minutes in the dark at RT at 500 rpm. After a further three washes, the plate was incubated in 50 µl of buffer A for 30 minutes. Plates received an additional wash and, after a final addition of 100 µl 1xPBS, were read using the Luminex MAGPIX® analyser. At least 50 beads per analyte were acquired per sample and MFI data were used for analysis.

This protocol is used to screen samples from regions that may be co-endemic for both schistosomiasis and malaria, potentially causing non-malaria reactivity against GST-tagged fusion proteins. Therefore, GST-coupled beads were included to quantify any GST-specific immunoglobulin (IgG) responses, which can be subtracted from total MFI against GST-tagged fusion proteins to better reflect malaria-specific IgG responses. Beads coupled with tetanus toxoid vaccine protein (TT) were also included as an internal assay quality control, given that measurable MFI values are expected for positive controls from regions where tetanus vaccine coverage is high. All data reported and analysed are in units of background subtracted MFI (Net MFI - blank wells).

Study samples used to develop standardisation methods
Samples used to validate data normalisation were based on all-age cross-sectional surveys conducted in July 2013 and
December 2013 in two villages in the West Coast Region and two villages in the Upper River Region (N=1,813) of The Gambia. Samples were eluted from a 6-mm dried blood spot (DBS) punch, corresponding to 4 μl of whole blood, and shaken overnight at room temperature in 200 μl of elution buffer containing 1xPBS, 0.05% sodium azide and 0.05% Tween-20, yielding an initial 1:50 sample dilution. At least 1 day prior to assay processing, samples were further diluted to a final 1:500 dilution using 10 μl of the 1:50 pre-dilution sample and 90 μl of blocking buffer B with E.coli extract to prevent non-specific binding. Negative and positive controls were also incubated one day prior in buffer B with E. coli, with negative controls prepared at a 1:500 dilution and Gambian pooled positive controls in a 6-point 5-fold serial dilution (1:10 – 1:31,250). Two wells on each plate containing only antigen-coupled beads and buffer B, but absent of any human serum, were included to measure background signal. A pool of 22 serum samples from malaria hyper-immune individuals in Upper River Region, The Gambia were used as a positive control, and plasma from 10 European malaria-naive adults were used as negative controls.

Estimating antigen-specific limits of quantification
To estimate the lower limit of quantification (LLOQ) and higher limit of quantification (HLOQ) for each antigen, two positive controls were tested (WHO NIBSC 10/198 and Tanzanian pooled positives (CP3)) using a 16-point 3-fold serial dilution, starting at 1:2. For each antigen and positive control, standard curves were fitted with Equation 1. The LLOQ MFI was defined as the MFI value where the upper 95%CI of the MFI parameter estimate is equivalent to the lower 95%CI of the standard curve estimate, and the HLOQ MFI where the lower 95%CI of the MFI parameter estimate is equivalent to the upper 95%CI of the standard curve estimate.

Quality control
For quality control of samples from The Gambia, Levey-Jennings charts were used to plot the mean MFI values of three concentrations from the positive control standard curve (high, 1:10; medium, 1:50; and low, 1:250) as well as the background values for each plate. The acceptable range of MFI values for inclusion in data analysis was defined as the mean ± two standard deviations of a subset of ten reference plates (selected based on the quality and consistency of their standard curve fits). Plates with MFI values outside this range for at least two standard curve dilutions and at least three antigens were rejected and repeated. Assays were processed in-country using beads transported from London. To assess the potential impact of interruption to the cold chain on bead stability, points on the Levey-Jennings plots were also ordered by date of plate processing and linear regression used to test for potential changes over time.

Immunoperoxidase normalisation
To account for observed between plate variation in positive control standard curves, data were adjusted using a loess normisation method. This method was tested using cross-sectional samples from The Gambia, as described above. First, positive control standard curves for each plate of antibody concentrations versus MFIs were fitted using a 4-parameter logistic equation:

$$MFI = MFI_{\text{max}} + \frac{(MFI_{\text{max}} - MFI_{\text{min}})}{1 + \left(\frac{\text{dilution}}{MFI_{\text{50}}^\text{dilution}}\right)^\text{slope}}. \quad (\text{Equation 1})$$

where $MFI_{\text{max}}$ is the upper asymptote or maximum MFI response of the standard curve, $MFI_{\text{min}}$ is the lower asymptote or minimum MFI response of the standard curve, $MFI_{\text{50}}$ is 50% of $MFI_{\text{max}}, \text{dilution}$ is positive control serum sample serial dilution, and $\text{slope}$ is the Hill coefficient or slope factor of the dose-response curve. $EC_{50}$ is the concentration or dilution that corresponds to $MFI_{\text{50}}$.

Next, ten reference plates from the study were selected based on the quality and consistency of their standard curve fits. For each antigen, a composite standard curve was computed by calculating the mean MFI values for the reference plates for 100 dilutions between the highest and lowest dilution on the standard curve. For each plate, the plate-to-reference standard curve MFI difference ($\Delta MFI$) was calculated for these 100 concentration points and a loess regression fit to $\Delta MFI$ as a function of mean MFI. The raw MFI data for all samples on the plate were then adjusted by the predicted $\Delta MFI$ based on the loess regression fit. Data were not corrected for background signal given that the between plate variation was already accounted for in the loess normalisation and all background MFIs were below 30 and therefore negligible.

Ethical approval
Written informed consent was obtained for all study participants. Ethical approval for the use of the Tanzanian samples was obtained from the institutional review boards of the National Institute of Medical Research of Tanzania, Kilimanjaro Christian Medical Centre (KCMC), and the London School of Hygiene & Tropical Medicine (LSHTM). For the Uganda (PRISM) samples, ethical approval was obtained from the Makerere University School of Medicine Research and Ethics Committee (REC REF 2011-203), Uganda National Council for Science and Technology (HS 1074), LSHTM Ethics Review Committee (Reference 6012) and the University of California, San Francisco on Human Research (Reference 027911). Finally, for the collection of the Gambian samples ethical approval was granted by the Scientific Coordinating Committee of the Medical Research Council (MRC) Laboratories in The Gambia and by the Joint MRC/Gambian Government Ethical Committee.

Results
Antigen to microsphere coupling
Optimal protein concentration for microsphere coupling, as determined by the titration with the MFI value closest to $EC_{50}$ of the dose-response curve, varied by antigen and ranged from as low as 0.017 ng/5000 beads for PfGLURP.R2 to 618 ng/5000 beads for GEXP18 (Figure 2, Table 3), depending on the immunogenicity of the recombinant protein. While some variation in
estimated EC\textsubscript{50} values for each antigen was observed between sample dilutions and positive controls (Supplementary Table S1), selecting the median EC\textsubscript{50} across all conditions as the optimal antigen-coupling concentration translated to MFI values on the linear portion of the dose-response curve, resulting in antibody responses measurable within a stable analytical range (Figure S1–Figure S3). A sigmoidal curve could be fit to the data for all antigens except Hyp2, where the midpoint between the two lowest titrations was selected as the optimal antigen concentration instead of the EC\textsubscript{50} (Table 3, Supplementary Table S1).

**Bead storage stability**

The effect of storage temperature on MFI signal over time varied by antigen, based on multivariate linear regression of change in MFI by storage week adjusted for temperature and serum sample dilution (Figure 3, Supplementary Table S2 and Supplementary Table S3). Compared to storage at −20°C, all tested temperatures (22°C, 37°C, and 42°C) were associated with a decrease of over 1,000 MFI per week of storage for GEXP18, HSP40.Ag1, Etramp4.Ag2 and Hyp2. For Etramp5.Ag1, PfMSP1\textsubscript{19} and PfGLURP.R2, degradation of more than 1,000 MFI per week was only observed at storage temperatures of 37°C and 42°C. No significant decreases in MFI signal were observed for PfAMA1 at any storage temperature. There was no significant degradation in signal between samples tested with beads stored for 6 months at 4°C compared to less than 4 weeks at 4°C, based on linear regression of logMFI with respect to time, after adjusting for antigen and allowing for random effects for sample and replicate.
Figure 3. Bead stability by temperature, storage time and dilution factor. Difference in the median fluorescence intensity (ΔMFI), of antigen-coupled beads stored at 22°C, 37°C and 42°C (compared to reference storage temperature of -20°C) after 1, 2, 4 and 8 weeks, and tested at six different positive control sample dilutions. Boxplots are based on data across all serum sample dilutions with median and interquartile range shown at each time point.
Buffer composition
In the absence of clarified *E. coli* lysate as a blocking agent against non-specific antibody binding to bacterial proteins, no significant differences in mean MFI were observed between the buffer compositions for both PfAMA1 and GEXP18 (Figure 4, Supplementary Table S4). The addition of the bacterial lysate was not associated with significant differences in MFI for yeast-produced PfAMA1. However, for GEXP18, the addition of the bacterial lysate resulted in a significant reduction in non-specific binding for both malaria endemic (-3,203.50 MFI, p=0.035) and malaria naïve samples (-5,857.30 MFI, p<0.001). The combination of buffer B and *E. coli* lysate did not have a synergistic reduction on background reactivity.

Serum sample dilution
Across all antigens tested, compared to serum dilution of 1:100, reductions in MFI values for negative controls were observed for sample serum dilutions of 1:500 (-911 MFI, p<0.001) and 1:1000 (-1,053, p < 0.001). These same serum dilutions maintained higher positive control values compared to a serum dilution of 1:2000 (Supplementary Figure S7). After adjusting for antigen, positive controls had an increased signal of 5,986 MFI (p<0.001) at a serum dilution at 1:500 and 3,006 MFI (p<0.001) at a serum dilution of 1:1000 compared to the average MFI signal at 1:2000. Higher sensitivity and specificity values were observed for serum dilutions of 1:500 (AUC 0.94 95%CI 0.91-0.96) and 1:1000 (AUC 0.96 95%CI 0.94-0.98) compared to serum dilutions of 1:100 (AUC 0.89 95%CI 0.86-0.93) and 1:2000 (AUC 0.91 95%CI 0.89-0.95), based on continuous MFI values across all antigens.

Limits of quantification
The limits of quantification of the assay differed by antigen and positive control used (Figure 5, Table 4). The highest HLOQs estimated were for PfAMA1, PfGLURP.R2, Etramp5.Ag1 and Etramp4.Ag2 (greater than 40,000 MFI using the CP3 Tanzanian positive control and greater than 30,000 MFI using the WHO reference standard). Estimated HLOQs for PfMSP1<sub>19</sub>, GEXP18, and HSP40.Ag1 ranged between 20,859 and 32,135 MFI across both positive controls, while the lowest estimated HLOQ was for Hyp2 (below 20,000 MFI for both controls). The lowest estimated LLOQs were for PfMSP1<sub>19</sub>, GEXP18, Hyp2, HSP40.Ag1 (below 500 MFI using the CP3 positive control), while estimated LLOQs for PfAMA1, PfGLURP.R2, Etramp5.Ag1, and Etramp4.Ag2 ranged 926–3,439 MFI across both positive controls tested.

Based on the CP3 positive control, the dynamic range was larger compared to the WHO reference control (e.g., both a lower LLOQ and higher HLOQ) for nearly all antigens including PfMSP1<sub>19</sub>, PfAMA1, Etramp5.Ag1, Etramp4.Ag2, GEXP18, and Hyp2. PfGLURP.R2 showed a higher HLOQ based on Tanzanian positive controls compared to the WHO reference standard, but also had a higher LLOQ. On the other hand, HSP40.Ag1 showed a lower LLOQ based on CP3 compared to the WHO reference, but also had a lower HLOQ.

Quality control
A total of 7,868 blood samples from The Gambia cross-sectional study were processed (96×96-well plates). Out of 96 plates, 3 fell outside the acceptable range of MFI values and were repeated. Degradation of MFI signal after exposure of antigen coupled beads to storage conditions above room temperature was also observed over a period of two months of sample processing, based on linear regression of MFI by date of plate processing (Supplementary Figure S8). For positive controls run at a dilution of 1:250, mean MFI for PfAMA1 at the start of sample processing was 24,480 and decreased at a rate of
Figure 5. Limits of quantification. Median fluorescence intensity (MFI) values are shown for pooled Tanzanian hyper-immune serum CP3 (blue) and WHO reference serum (red) in a 12-point serial dilution. Horizontal lines represent the mean and 95% confidence intervals of the upper and lower asymptotes of the sigmoidal curve for CP3 (blue), WHO (red), and the mean background MFI in green.
Table 4. Limits of quantification at 1:400 serum sample dilution. Units in median fluorescence intensity.

<table>
<thead>
<tr>
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<th>CP3</th>
<th>WHO</th>
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<td>HLOQ</td>
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<td>52,967</td>
</tr>
<tr>
<td>Hyp2</td>
<td>132</td>
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</tr>
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61 MFI per plate (p=0.0104) over the course of two months. For GEXP18, mean MFI was 8,209 at the start of sample processing and declined at a rate of 31 MFI per plate (p=0.007) over the same period.

Data normalisation

Proportional differences in plate-specific MFI values compared with mean MFI values of reference plates (indicated as ΔMFI in Figure 6) were highly dependent on the MFI range (Figure 6, Supplementary Figures S9-11). In other words, MFIs in the higher end of responses could show larger between-plate variations than MFIs in the lower end of responses (or vice versa) and may not be easily adjusted for using one proportional factor across the full range of MFI values—the method typically used for normalising ELISA-based data. The extent of the variations differed by plate and antigen. Therefore, the loess normalisation allowed for raw data to be adjusted.

Figure 6. Loess normalisation for PfAMA1 and GEXP18. Loess normalisation of antigens PfAMA1 and GEXP18 is illustrated for one example plate based on cross-sectional samples from The Gambia. Panels A and B show the loess fit (red) and linear fit (blue) of ΔMFI on the y-axis (the difference in unadjusted MFI of the standard curve of a single plate and the mean MFI of all standard curves from ten reference plates) and mean MFI on the x-axis. Panels C and D show the raw MFI values of individual samples on the x-axis versus normalised MFI values on the y-axis, and the equality line is shown diagonally in red.
and weighted according to MFI range-specific differences (Figure 6, Supplementary Figures S9-11). Additionally, loess regression provided a better fit to the ΔMFI versus mean MFI data compared to linear regression, indicating that it may provide better adjustment values for plate-specific normalisation.

Discussion

This study builds on previous work optimising the Luminex® qSAT as a multiplex platform of *Plasmodium* antigens. To develop a panel of serological markers for the characterisation of both historical and recent malaria exposure, the methods tested here aimed to standardise a protocol that could consistently measure a large range of antibody responses. This included the optimisation of antigen-to-bead coupling concentrations, testing of coupled-bead stability at a range of temperatures reflective of variable storage conditions across laboratories, as well as buffer composition and serum sample dilution to minimise non-malaria specific background reactivity.

Results showed that storage of antigen-coupled beads at temperatures below 37°C, and ideally at room temperature or lower, minimised degradation of MFI signal over an 8-week period for all antigens. Some antigens, such as *Pf*AMA1, exhibited more stability than others, where no significant degradation in MFI signal was observed over an eight-week period even at temperatures up to 42°C. Storage at room temperature also affected the bead stability for GEXP18, HSP40.Ag1, Etramp4, Ag2 and Hyp2. This is consistent with previous studies reporting that stability of beads is antigen-specific. This suggests that additional formulations to improve bead stability, such as lyophilisation in the presence of stabilisers, could be explored to minimise the impact on assay results. This is particularly critical when reagents are subject to transport or storage conditions with a high risk of interruption to the transport cold chain. Temperatures of 22°C and above were tested in this study to assess the potential impact of disrupted cold chain during shipping to or long-term storage in laboratories in low- and middle-income settings. Further work should also investigate bead stability during long-term storage at 4°C, the standard storage condition for most research groups using Luminex platforms.

Optimal antigen-to-bead coupling concentrations were determined for eight *Pf* antigens, including five markers of recent malaria exposure1. These concentrations were selected to cover a range of positive controls and serum sample dilutions allowing for generalisability across multiple study conditions. Previous studies have found that bead coupling concentration is the most consistent factor influencing assay variability. Failing to control the density of antigens on the microsphere surface may result in either sub-optimal coating and low reactivity or over-coating and subsequent precipitation or aggregation of beads on the bottom of the plate, impairing surface suspension antibody binding1. The degree to which this occurs and the tendency for non-uniform protein aggregation may also be antigen-specific, depending on the physical characteristics of protein structure. New coupling methods are currently in development, including biotinylated microspheres that allow coupling to specific antigen regions for more consistent protein orientation. Future work should validate the analytical reproducibility of these coupling concentrations by monitoring the consistency in MFI signal between bead sets and recombinant protein batches. This may also identify antigens that may be better suited on either bead-based or microarray platforms. Given that protocols are being developed for high-throughput processing, the selection of optimal bead coupling concentrations should ideally ensure a consistent analytical signal across bead sets (i.e., close to the EC₅₀ point on the standard curve), while minimising the amount of antigen needed to achieve cost and volume efficiencies over time. If these analytical and operational challenges can be met across a diverse set of antigens, there may be strong potential for including these analytes on larger multi-disease Luminex panels.

The inclusion of *E. coli* lysate in the sample incubation buffer was found to be an effective blocking agent, particularly against non-specific antibody binding to antigens expressed in *E. coli* by significantly reducing the MFI signal in negative samples. The impact of the addition of the *E. coli* lysate is suggestive of co-purification of *E. coli* proteins, for which there was little evidence by ELISA but are likely only detected on the Luminex platform due to the higher dynamic range. Moving forward, the inclusion of additional chromatography separation techniques would minimise the presence of any co-purifying proteins and potentially negate the need for the *E. coli* lysate additive. Serum sample concentration was also optimal at both 1:500 and 1:1000 to reduce non-specific background reactivity in negative samples while retaining a measurable degree of reactivity in positive samples. These serum dilutions had higher sensitivity and specificity for identifying positive control sera compared to dilutions of 1:100 and 1:2000. Several other assay conditions have been validated in previous studies. This includes the testing of 1,000 compared to 2,000 beads per well13,14, the use of plasma compared to DBS15, and the comparison of single-plex with multiplex platforms16,17. Based on these studies, the protocol presented here uses 1,000 beads as an optimal baseline condition. Additionally, processing samples in duplicate, the use of plasma, and single-plex platforms did not significantly improve assay results. Therefore, optimal conditions confirmed from previous studies were incorporated into this protocol but not investigated further.

This study also reports antigen-specific limits of quantification (LLOQs and HLOQs) for several new *Pf* recombinant proteins, where the dynamic range varied between antigens and positive controls used. The WHO reference standard was found to have a narrower dynamic range compared to in-house positive control sera, which has also been observed in previous studies using the WHO positive control18. This is likely due to the selection of sera for this reference standard based on small number of antigens primarily associated with long-lived antibody responses (*Pf*MSPI19, *Pf*AMA1, and *Pf*CSP) and from only one geographical location (Kenya). Therefore, it may not be well suited for capturing short-lived antibody responses in populations from other endemic settings. Future work should consider how reference standards can be further improved to maximise the
measurable range for a larger selection of Pf antigens and across different geographical regions and also determine the LOQs across a range of sample serum dilutions currently being used.

Several challenges remain for the further development of multiplex serological platforms for malaria surveillance. First, many Luminex-based malaria studies currently use antibody concentrations estimated from the positive control standard curve for epidemiological analysis. As already reported in previous studies, assay conditions can affect the fit of the standard curve used to normalise the data to arbitrary concentration units. This potentially results in large deviations in concentration estimates. MFI responses measured independently from a standard curve might reflect true variation, while normalisation methods that convert to relative concentration values will be more influenced by the precision of the statistical fit of the standard curve. Therefore, we use a loess normalisation method for standardising data between assays, which attempts to maintain units in MFI values while also accounting for dilution-dependent variability in signal (i.e., magnitude of between plate variation that differs by MFI range). The robustness of this procedure, however, should be validated with additional laboratory data. For example, repeat testing of one set of endemic sera across multiple plates to simulate assay variability can be used to estimate residual between-plate variance after data normalisation and to determine if the variance is antigen specific. An assessment of potential overfitting using loess compared to linear regression or other normalisation methods should also be explored. It is also important to note that because normalisation is dependent on the standard curve, and adjustment of data below or above the minimum and maximum MFI values of the positive control titrations should be avoided as it may be difficult to extrapolate the relationship at these extremes (as seen with PmMSP119, Supplementary Figure S9). Therefore, designing standard curves to include titrations matching the range of MFI values expected in the samples being processed is important when using this normalisation method.

The ultimate utility of these assays will be the application to high-throughput processing in endemic country laboratories. Minimal specialist equipment is required aside from a Luminex reader, a magnetic plate, and coupled microspheres and standard ELISA laboratory reagents. Therefore, Luminex-based processing is feasible in a range of endemics settings, from local hospitals with back-up electric generators (with consistent supply for at least two 90 minutes incubations and a one hour plate reading) to larger laboratories in urban settings. This will likely require further validation of bead stability during field-based transport and storage conditions. To confirm the reproducibility of the assay and data standardisation methods, between laboratory and user variability testing should be explored. The application of these data to epidemiological analysis will also need to consider antigen-specific kinetics across different endemic settings, in order to validate the use of novel serological markers of recent malaria exposure for routine surveillance or the evaluation of community-based efficacy trials. Standardised methods for determining sero-positivity thresholds for new antigens not previously used should be established.

Data availability
Underlying data
Raw data for this study, including output data for antigen and serum sensitivity and stability testing, output data from qSAT assays and data for validation and standardisation, are available on OSF. DOI: https://doi.org/10.17605/OSF.IO/AUJ35.

Extended data
Extended data are available on OSF. DOI: https://doi.org/10.17605/OSF.IO/AUJ35.

Figure S1. Titration of antigen-concentration for bead coupling, across five serum sample dilutions and two positive controls for PfMSP1.19, PfAMA1, and PGLURP.R2. Filled circles represent the EC50 specific to the serum dilution and positive control. Vertical black line is the median EC50 concentration across all serum dilutions and positive controls.

Figure S2. Titration of antigen-concentration for bead coupling, across five serum sample dilutions and two positive controls for Etramp5.Ag1 and Etramp4.Ag2. Filled circles represent the EC50 specific to the serum dilution and positive control. Vertical black line is the median EC50 concentration across all serum dilutions and positive controls.

Figure S3. Titration of antigen-concentration for bead coupling, across five serum sample dilutions and two positive controls for GEXP18 and HSP40.Ag1. Filled circles represent the EC50 specific to the serum dilution and positive control. Vertical black line is the median EC50 concentration across all serum dilutions and positive controls.

Figure S4. Median fluorescence intensity (MFI) for PfMSP1 and PGLURP.R2 of positive and negative samples for each buffer composition. Buffer compositions tested include buffer A (red), buffer A with *E. coli* lysate (pink), buffer B (blue), and buffer B with *E. coli* lysate (light blue). MFI for positive samples shown in colour (left) and corresponding negatives samples in grey (right) for each buffer composition.

Figure S5. Median fluorescence intensity (MFI) for Etramp5.Ag1 and Etramp4.Ag2 of positive and negative samples for each buffer composition. Buffer compositions tested include buffer A (red), buffer A with *E. coli* lysate (pink), buffer B (blue), and buffer B with *E. coli* lysate (light blue). MFI for positive samples shown in colour (left) and corresponding negatives samples in grey (right) for each buffer composition.

Figure S6. Median fluorescence intensity (MFI) for HSP40.Ag1 and Hyp2 of positive and negative samples for each buffer composition. Buffer compositions tested include buffer...
A (red), buffer A with *E. coli* lysate (pink), buffer B (blue), and buffer B with *E. coli* lysate (light blue). MFI for positive samples shown in colour (left) and corresponding negatives in grey (right) for each buffer composition.

**Figure S7.** Serum sample dilution optimisation. Mean median fluorescence intensity (MFI) of positive and negative samples tested at four serum sample dilutions (1:100, pink; 1:500, blue; 1:1000, green; 1:2000, purple). Median MFI of negative samples are shown in grey to the right of positive samples shown in colour.

**Figure S8.** Levey-Jennings plots for Luminex plate quality control. Solid points represent the median fluorescence intensity (MFI) values of positive controls, ordered left to right by date of plate processing. Solid horizontal lines represent MFI values of either one or two standard deviations from the mean. Coloured lines are the linear regression fit (mean and 95% CI) of change in MFI by date of plate processing, representing estimated signal degradation over a period of 2 months.

**Figure S9.** Loess normalisation *P/MSP1* and *P/GLURP.R2*. Loess normalisation of antigens *P/MSP1.19* and *P/GLURP.R2* is illustrated for one example plate based on cross-sectional samples from The Gambia. Upper two panels show the loess fit (red) and linear fit (blue) of Δ MFI on the y-axis (the difference in unadjusted MFI of the standard curve of a single plate and the mean MFI of all standard curves from ten reference plates) and mean MFI on the x-axis. Lower two panels show the raw MFI values of individual samples on the x-axis versus normalised MFI values on the y-axis, and the equality line is shown diagonally in red.

**Figure S10.** Loess normalisation Etramp5.Ag1 and Etramp4.Ag2. Loess normalisation of antigens Etramp5.Ag1 and Etramp4.Ag2 is illustrated for one example plate based on cross-sectional samples from The Gambia. Upper two panels show the loess fit (red) and linear fit (blue) of Δ MFI on the y-axis (the difference in unadjusted MFI of the standard curve of a single plate and the mean MFI of all standard curves from ten reference plates) and mean MFI on the x-axis. Lower two panels show the raw MFI values of individual samples on the x-axis versus normalised MFI values on the y-axis, and the equality line is shown diagonally in red.

**Figure S11.** Loess normalisation HSP40.Ag1 and Hyp2. Loess normalisation of antigens HSP40.Ag1 and Hyp2 is illustrated for one example plate based on cross-sectional samples from The Gambia. Upper two panels show the loess fit (red) and linear fit (blue) of Δ MFI on the y-axis (the difference in unadjusted MFI of the standard curve of a single plate and the mean MFI of all standard curves from ten reference plates) and mean MFI on the x-axis. Lower two panels show the raw MFI values of individual samples on the x-axis versus normalised MFI values on the y-axis, and the equality line is shown diagonally in red.

### References


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Version 2

Reviewer Report 27 April 2020

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Takafumi Tsuboi
Division of Malaria Research, Proteo-Science Center, Ehime University, Matsuyama, Ehime 790-8577, Japan

All of my comments have been appropriately addressed.

Competing Interests: No competing interests were disclosed.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Version 1

Reviewer Report 16 May 2019

https://doi.org/10.21956/wellcomeopenres.16307.r35431

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? Takafumi Tsuboi
Division of Malaria Research, Proteo-Science Center, Ehime University, Matsuyama, Ehime 790-8577, Japan

Antibody responses have been used to characterize transmission and exposure history in malaria-endemic settings for over a decade. The authors are one of the leading groups on such studies, especially P. falciparum seroepidemiology. In this study, they carefully validated their Luminex-based antibody measurement system using eight P. falciparum recombinant antigens, associated with long- and
short-lived antibody responses. They finally standardized antigen-specific bead
coupling concentrations, buffer compositions, storage conditions, limits of quantifications and
normalization strategies. The results in this work will be very useful for the current and future expansion of
the inclusion of new antigens in this Luminex-based platform. I have a few comments that need to be
considered:

1. For which field setting can the MAGPIX platform be applicable (Target Product Profile)? For example, a
local hospital with back-up electric generator or a bigger laboratory in the central city.

2. 6 out of 8 antigens used in this study were expressed as GST-fused recombinant proteins in E. coli as
shown in Table 1. GST and TT were also included as negative controls. 1) Please clarify how to use MFI
on negative controls to normalize MFI on GST-fused antigens. 2) I have another concern about using
GST-fused recombinant proteins as antigens for serology because there are many endemic areas of both
Schistosomiasis and malaria. If the people had a history or co-infection with this worm, the interpretation
of the results will be very difficult. Do the authors have any idea how to overcome this challenge?
Therefore, I strongly recommend the authors to use another purification tag such as His-tag in the future
development.

Is the rationale for developing the new method (or application) clearly explained?
Yes

Is the description of the method technically sound?
Yes

Are sufficient details provided to allow replication of the method development and its use by
others?
Partly

If any results are presented, are all the source data underlying the results available to ensure full
reproducibility?
Yes

Are the conclusions about the method and its performance adequately supported by the
findings presented in the article?
Yes

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Malaria protein research, Malaria vaccine research, Malaria molecular biology

I confirm that I have read this submission and believe that I have an appropriate level of
expertise to confirm that it is of an acceptable scientific standard, however I have significant
reservations, as outlined above.

Author Response 07 Apr 2020
Kevin Tetteh, London School of Hygiene & Tropical Medicine, London, UK
Reviewer 2

We would like to thank Dr. Tsuboi for his useful comments.

For which field setting can the MAGPIX platform be applicable (Target Product Profile)? For example, a local hospital with back-up electric generator or a bigger laboratory in the central city.

Together with our international colleagues, we have demonstrated the utility of the MagPix platform in a number of different settings as evidenced in the following publications (Surendra H et al. BMC Med. 2020 Jan 28;18(1):9. doi: 10.1186/s12916-019-1482-7. [PMID: 31987052]; van den Hoogen Sci Rep. 2020 Jan 24;10(1):1135. doi: 10.1038/s41598-020-57876-0. [PMID: 31980693]; Achan J et al. Clin Infect Dis. 2019 Aug 12. pii: ciz740. doi: 10.1093/cid/ciz740. [PMID: 31980693]). We and others have shown that the platform can be used in a wide range of study sites as long as there is access to an uninterrupted power supply. Apart from the Magpix reader, magnetic plate and coupled microspheres, no additional specialist equipment is required, outside of typical laboratory materials. “Minimal specialist equipment is required aside from a Luminex reader, a magnetic plate, and coupled microspheres and standard ELISA laboratory reagents. Therefore, Luminex-based processing is feasible in a range of endemics settings, from local hospitals with back-up electric generators (with consistent supply for at least two 90 minutes incubations and a one hour plate reading) to larger laboratories in urban settings.”

6 out of 8 antigens used in this study were expressed as GST-fused recombinant proteins in *E. coli* as shown in Table 1. GST and TT were also included as negative controls. 1) Please clarify how to use MFI on negative controls to normalize MFI on GST-fused antigens. 2) I have another concern about using GST-fused recombinant proteins as antigens for serology because there are many endemic areas of both Schistosomiasis and malaria. If the people had a history or co-infection with this worm, the interpretation of the results will be very difficult. Do the authors have any idea how to overcome this challenge? Therefore, I strongly recommend the authors to use another purification tag such as His-tag in the future development.

- Tetanus toxoid (TT) and GST are not intended as ‘negative controls’. TT was included as an internal positive control as it is expected that most samples, including malaria endemic and naïve controls, will have measurable MFI values against the TT coupled beads and therefore confirm that the assay was run successfully. GST only responses are measured and subtracted from target responses to account for any possible non-specific GST reactivity in samples.
- We understand the reviewer’s concern. The GST-tag is based on a glutathione s-transferase from *Schistosoma mansoni*, which is prevalent in some of the malaria endemic sites under test. However, we and others have found that subtracting the non-specific, or potentially cross-reactive responses to the fusion tag has been demonstrated elsewhere to be effect in measuring antigen specific responses (Dobaño C et al J Infect Dis 2008 [PMID: 18260767]; Ahlborg N et al Clin Exp Immunol 2002 [PMID: 12165089]). That said, we are currently exploring other expression platforms including his-tag to eliminate this concern in the future.

**Competing Interests:** No competing interests were disclosed.
The authors describe comparisons of immunoassay conditions utilizing a bead-based system as applied to a *Plasmodium falciparum* 8-antigen panel. Data is provided for how antigen coupling concentrations are selected, differences in sample dilution and blocking buffers, and coupled bead storage conditions. Additionally, the authors outline a normalization scheme when multiple assay plates are run for a study. The report is timely, as multiple malaria groups are now using bead-based multiplex assays and investigating new antigens to augment panels. The authors could provide some additional information regarding the details of their assay and reporting, as well as make some text in the manuscript clearer for the inexperienced reader. Furthermore, some figures currently show data for all 8 Pf antigens, whereas others only display data for 2 or 3. At the least, the data for all 8 antigens should be included as Supplementary data.

**Minor suggestions:**

1. Are references available for more detailed information of recombinant antigen production for the 8 presented here (or similar)?

2. Explain in Methods how the beads were washed.

3. Provide detail about GST and tetanus toxoid couplings. Explain what is meant by: “to allow for correction of any GST-specific immunoglobulin”.

4. Explain how “clarified *E. coli* lysate” was produced.

5. Are MFI values reported or MFI-background values?

6. The current flowchart in Figure 1 is a little difficult to read, especially for someone unfamiliar with this assay. It is suggested to add some more text to the boxes to clarify and read more like a protocol. For example, one box reads: “8 uL coupled microspheres, 5 mL buffer A, (1,000 beads per well)”, and would be clearer with: “8 uL of coupled beads added to 5 mL buffer A for a concentration of approximately 1,000 beads per well”.

7. Titration curves in Figure 2 for the other antigens would be helpful in Supplementary.

8. Unfortunately, in Figure 3, the authors do not include a storage condition of 4 degrees which most researchers will choose to keep their coupled beads at for practical storage. It’s not clear why conditions starting at room temperature and increasing were chosen, as this wouldn’t be utilized by a group having MAGPIX technology. Even an interruption of cold-chain would only be for a short
period of time for this type of work.

9. For the Buffer Composition comparisons, the authors only offer data for two Pf antigens in Figure 4 and Supplementary Table 4. Please provide this same comparison for all 8 Pf antigens as well as the two control proteins. The same for Figure 6 in that all antigens should at least be shown in Supplementary.

Is the rationale for developing the new method (or application) clearly explained?
Yes

Is the description of the method technically sound?
Yes

Are sufficient details provided to allow replication of the method development and its use by others?
Partly

If any results are presented, are all the source data underlying the results available to ensure full reproducibility?
Yes

Are the conclusions about the method and its performance adequately supported by the findings presented in the article?
Yes

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Malaria serological assays

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

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Author Response 07 Apr 2020

Kevin Tetteh, London School of Hygiene & Tropical Medicine, London, UK

Reviewer 1
We would like to thank the reviewer for their useful comments:

Are references available for more detailed information of recombinant antigen production for the 8 presented here (or similar)?

reflect this:

“Each antigen was generated and expressed in Escherichia coli as glutathione S-transferase (GST)-tagged fusion proteins using methods as described elsewhere (PMID: 29902183, PMID: 23897617, PMID: 12654798). The exception to the panel was PfAMA1, which was expressed in Pichia pastoris as a histidine-tagged protein (PMID: 17192270).”

**Explain in Methods how the beads were washed.**

The following sentences have been included into the qSAT procedure to clarify the wash step.

“Next, 50 μl of this combined microsphere mixture was added to a 96-well flat bottom plate (BioPlex Pro™, Bio-Rad Laboratories, UK) and washed once by placing the assay plate onto a magnetic plate separator (Bio-Plex®, Bio-Rad Laboratories, UK) and incubated for 2 minutes at room temperature. Plates were then inverted forcefully to remove the liquid and 100 μl of PBS-T (1xPBS, 0.05% Tween-20) added to each well. “

**Provide detail about GST and tetanus toxoid couplings. Explain what is meant by: “to allow for correction of any GST-specific immunoglobulin”**

Tetanus toxoid (TT) coupled beads were included as the Tetanus vaccine is widely used and therefore useful as an internal control. It would be expected that most samples, including malaria endemic and naïve controls will have measurable MFI values against the TT coupled beads and therefore confirm that the assay was run successfully. This is a particularly useful control when samples are assayed as single wells (with no duplicate result to compare to).

The GST-tag used in the purification of the recombinant proteins is based on a glutathione s-transferase from Schistosoma mansoni, a blood trematode common to some of the malaria endemic sites under test. As several of the antigens are expressed as GST-tagged fusion proteins it is important to correct for any background, or non-malaria specific reactivity to the GST-tag. Assaying for reactivity to beads coupled with purified GST allows us to measure the level of reactivity to the tag, if any, and background correct the response by subtracting the GST response. Subtraction of non-specific, or potentially cross-reactive responses to fusion tags in serological studies has been demonstrated elsewhere (Dobaño C et al J Infect Dis 2008 [PMID: 18260767]; Ahlborg N et al Clin Exp Immunol 2002 [PMID: 12165089])

**Explain how “clarified E. coli lysate” was produced.**

To produce the bacterial lysate, untransformed E.coli were cultured using the Studier recipe for autoinduction media under the same conditions as used for expression of the recombinant protein with one exception. The addition of an antibiotic for selection of recombinants is omitted. After culturing for approximately 18 hours at 37°C, the cells are harvested by centrifugation, resuspended in PBS and lysed using a high pressure homogeniser. The lysed material is then centrifuged at 16,000 rpm for 15 min to pellet the lysed E.coli membranes. The clarified lysate is then aliquoted and stored at -20°C until needed.

As sentence has been included in the text to summarise this point:

“Bacterial lysate was generated from the culture of untransformed E.coli and used in the preparation of assay buffers as non-specific protein to eliminate background reactivity to E.coli proteins (PMID: 26216993).”
Are MFI values reported or MFI-background values?

All results are reported as MFI-background and the following sentence has been added to the methods: “All data reported and analysed are in units of background subtracted MFI.”

The current flowchart in Figure 1 is a little difficult to read, especially for someone unfamiliar with this assay. It is suggested to add some more text to the boxes to clarify and read more like a protocol. For example, one box reads: “8 uL coupled microspheres, 5 mL buffer A, (1,000 beads per well)”, and would be clearer with: “8 uL of coupled beads added to 5 mL buffer A (for a concentration of approximately 1,000 beads per well)”.

We appreciate the reviewer’s comments and the flowchart has been edited accordingly and a new version uploaded.

Titration curves in Figure 2 for the other antigens would be helpful in Supplementary.

The titrations curves for the remaining antigens were previously included in the supplement (Figures S1, S2, and S3)

Unfortunately, in Figure 3, the authors do not include a storage condition of 4 degrees which most researchers will choose to keep their coupled beads at for practical storage. It’s not clear why conditions starting at room temperature and increasing were chosen, as this wouldn’t be utilized by a group having MAGPIX technology. Even an interruption of cold-chain would only be for a short period of time for this type of work

Coupled and uncoupled beads are stored at 4°C as standard. However, coupled beads are routinely shipped globally to collaborator sites, where disruptions to the cold chain can and has occurred. As such we felt it was necessary to evaluate the potential impact of suboptimal shipping and/or storage conditions on the functionality of the coupled beads. The following text has been included in the discussion as a point to consider in future studies: “Temperatures of 22°C and above were tested in this study to assess the potential impact of disrupted cold chain during shipping to or long-term storage in laboratories in low- and middle-income settings. Further work should also investigate bead stability during long-term storage at 4°C, the standard storage condition for most research groups using Luminex platforms.”

For the Buffer Composition comparisons, the authors only offer data for two Pf antigens in Figure 4 and Supplementary Table 4. Please provide this same comparison for all 8 Pf antigens as well as the two control proteins. The same for Figure 6 in that all antigens should at least be shown in Supplementary.

This has now been included in Supplementary data as Figures S4-6 for buffer testing, and Figures S9-11 for loess normalisation

**Competing Interests:** No competing interests were disclosed.
Comments on this article

Reviewer Response 25 Apr 2020

Takafumi Tsuboi, Ehime University, Matsuyama, Japan

All of my comments have been appropriately addressed.

**Competing Interests:** No competing interests were disclosed.