Pregnant women constitute a promising sentinel group for continuous monitoring of malaria transmission. To identify antibody signatures of recent *Plasmodium falciparum* exposure during pregnancy, we dissected IgG responses against VAR2CSA, the parasite antigen that mediates placental sequestration. We used a multiplex peptide-based suspension array in 2,354 samples from pregnant women from Mozambique, Benin, Kenya, Gabon, Tanzania, and Spain. Two VAR2CSA peptides of limited polymorphism were immunogenic and targeted by IgG responses readily boosted during infection and with estimated half-lives of <2 years. Seroprevalence against these peptides reflected declines and rebounds of transmission in southern Mozambique during 2004–2012, reduced exposure associated with use of preventive measures during pregnancy, and local clusters of transmission that were missed by detection of *P. falciparum* infections. These data suggest that VAR2CSA serology can provide a useful adjunct for the fine-scale estimation of the malaria burden among pregnant women over time and space.

A agile malaria surveillance and response systems that can be sustained over time are needed for the optimal design of control programs (1,2). Rates of *Plasmodium falciparum* infection among pregnant women are sensitive to changes in transmission (3,4) and correlate well with infection in infants (5) and children (6,7). Thus, passive detection of malaria cases at maternal health care services constitutes a promising approach to providing contemporary data on the levels, and changes in levels, of malaria burden in the population for successful malaria control and elimination (8).

After exposure to *P. falciparum* parasites that sequester in the placenta (9), antibodies against VAR2CSA, a multidomain variant antigen of the *P. falciparum* erythrocyte membrane protein 1 family, develop in pregnant women (10). VAR2CSA is expressed on the surface of infected erythrocytes and mediates placental sequestration of parasites through binding to chondroitin sulfate A (11). Levels of antibodies against VAR2CSA are affected by variables that influence the risk for *P. falciparum* exposure (12–14) and mirror malaria trends during pregnancy (3). Moreover,
levels of VAR2CSA IgG at delivery correlate with the risk for malaria in the offspring (14), suggesting the value of these antibodies for pinpointing areas of high malaria transmission (15). Because VAR2CSA antibodies persist after the infection is cleared (16), they can provide a sensitive adjunct for \textit{P. falciparum} monitoring, especially in areas of low malaria endemicity, where the chances of detecting antibodies are higher than those of detecting the parasite (17).

The utility of serosurveillance depends mainly on specific properties of the antigen, including immunogenicity, polymorphism, cross-reactivity, and longevity of the antibodies. Because different VAR2CSA domains elicit IgG responses with varying magnitudes and dynamics (16,18,19), we hypothesized that short-lived antibodies against immunogenic nonpolymorphic VAR2CSA epitopes would enable a fine-scale estimation of recent \textit{P. falciparum} transmission during pregnancy (17). We examined plasma from pregnant women living in areas in which \textit{P. falciparum} transmission varied from high to low and absent (Benin, Gabon, Mozambique, Kenya, Tanzania, and Spain) against a quantitative suspension array containing VAR2CSA and general parasite antigens. We first selected IgG responses that were rapidly acquired after \textit{P. falciparum} infection, did persist in circulation, and were sensitive to the level of parasite exposure in pregnant women from Mozambique and Spain. We then used the serologic assay to quantify the relationship of VAR2CSA antibody responses with \textit{P. falciparum} infection as well as with temporal, spatial, and intervention-driven changes in malaria burden among pregnant women.

\section*{Methods}

\subsection*{Study Sites, Population, and Procedures}

We included in our study pregnant women who participated in 3 clinical trials of intermittent preventive treatment during pregnancy (IPTp) during 2003–2005 in Mozambique (NCT00209781) (20) and during 2010–2012 in Mozambique, Benin, Gabon, Kenya, and Tanzania (NCT00811421) (21,22). Participants were recruited at their first antenatal visit, and all received a long-lasting insecticide-treated bed net. During 2003–2005, all received 2 doses of sulfadoxine/pyrimethamine (20); during 2010–2012, they received 2 doses of mefloquine or sulfadoxine/pyrimethamine if they were not HIV infected (21) and 3 doses of mefloquine or placebo plus daily cotrimoxazole prophylaxis if they were HIV infected (22). At delivery, tissue samples from the maternal side of the placenta, as well as 50 \textmu L peripheral and placental dried blood spots (DBS), were collected. Peripheral and placental blood from pregnant women in Mozambique and Benin were also collected into EDTA Vacutainer tubes (Becton Dickinson, https://www.bd.com) and centrifuged; plasma was stored at \textdegree 20\textcelsius. From a subset of pregnant women in Mozambique who delivered during 2011–2012, peripheral blood samples were also collected at the first antenatal visit and before administration of the second IPTp dose. We geocoded the households of women in Mozambique by using a global information system. Clinical malaria episodes were treated according to national guidelines at the time of the study (20–22). DBS and plasma samples were also collected from 49 pregnant women never exposed to \textit{P. falciparum} who delivered in 2010 at the Hospital Clinic of Barcelona (Barcelona, Spain).

The study was approved by the ethics committees from the Hospital Clinic of Barcelona, the Comité Consultatif de Déontologie et d’Éthique from the Institut de Recherche pour le Développement (Marseille, France), the Centers for Disease Control and Prevention (Atlanta, GA, USA), and national ethics review committees from each malaria-endemic country participating in the study. Written informed consent, which included permission to test for immune markers by using stored biological samples, was obtained from all participants.

\subsection*{Laboratory Determinations}

At recruitment, we assessed HIV serostatus by using rapid diagnostic tests according to national guidelines and hemoglobin level at delivery by using following mobile devices on capillary blood samples: HemoCue (Danaher, http://www.hemocue.com), Hemocontrol (EKF Diagnostics, http://www.ekfdiagnostics.com), and KX analyzer (Sysmex, http://www.sysmex.com). Thick and thin blood films and placental biopsy samples were checked for \textit{Plasmodium} spp. according to standard, quality-controlled procedures (3). We tested blood on filter paper for the presence of \textit{P. falciparum} in duplicate by means of a real-time quantitative PCR (qPCR) targeting 18S ribosomal DNA (3).

\subsection*{Antibody Measurements}

We measured IgG in plasma (Benin and Mozambique) or on DBS (Gabon, Kenya, and Tanzania) in appropriate conditions for plasma elution (19) by using the xMAP technology and the Luminex 100/200 System (https://www.luminexcorp.com) for 37% of pregnant women participating in the clinical trials with samples available. We constructed 2 multiplex suspension array panels (Appendix 1, https://wwwnc.cdc.gov/EID/article/25/10/18-1177-App1.pdf) (19), 1 including \textit{P. falciparum} recombinant proteins (VAR2CSA Duffy binding-like recombinant domains DBL3X, DBL5E, and DBL6E; apical membrane antigen 1 [AMA1]; and 19-kDa fragment of the merozoite surface protein-1 [MSP1\textsubscript{19}], from 3D7 strain) and 1 consisting of synthetic peptides (25 VAR2CSA peptides covering conserved and semiconserved regions of VAR2CSA and a circumsporozoite peptide [pCSP]) (19). To assess unspecific
IgG recognition, we used bovine serum albumin in both arrays (19). Procedures for reconstitution of DBS and quality control, bead-based immunoassay, data normalization, and definition of seropositivity cutoffs are described in Appendix 1.

var2csa Sequencing and 3D Protein Modeling
We used DNA extracted from 50 DBS that were P. falciparum positive by qPCR for Sanger sequencing of var2csa PCR amplification products covering peptides of interest (Appendix 1). Sequence variability with respect to the peptide included in the array was assessed after amino acid alignment, and a 3D model of the DBL1X-ID1 region was developed by using Chimera version 1.5.3 (https://www.cgl.ucsf.edu; Appendix 1).

Definitions and Statistical Analyses
We included in the analysis pregnant women for whom all information was available for IPTp, date of delivery, HIV status, age, parity, and antibody responses. We classified women as primigravid (first pregnancy) or multigravid (≥1 previous pregnancy) and categorized age as <20, 20–24, or ≥25 years (13). Anemia was defined as hemoglobin level at delivery <11 mg/L. We compared proportions by using the Fisher exact test. We used univariate regression models to evaluate the association of log-transformed IgG levels (linear) and seropositivity (logistic) with study periods (2004–2005 and 2010–2012) and country, P. falciparum infection, parity, anemia, and IPTp intervention, taking into account potential confounding variables (HIV and age) in multivariate models. We assessed the modification of the associations by HIV infection or parity by including interaction terms into the regression models. To control the false discovery rate in the selection of antigens, we computed adjusted p values (q-values) by using the Simes procedure (23). We used multilevel mixed-effect linear regression analysis to estimate half-life and time to double (T½) IgG levels in the longitudinal cohort of pregnant women from Mozambique (Appendix 1). We identified spatial clusters of P. falciparum infection and seropositivity as well as the most likely hotspots by using the Ward hierarchical cluster analysis and Kulldorff spatial scan method (Appendix 1). We performed statistical analyses by using Stata/SE software version 12.0 (StataCorp, https://www.stata.com), R statistics software version 3.2.1 (https://www.r-project.org), and Graphpad Prism version 6 (https://www.graphpad.com).

Results
Study Participants and P. falciparum Prevalence
Study participants consisted of 2,354 pregnant women (Table; Appendix 1 Figure 2) recruited during 2004–2005 (n = 146) and 2010–2012 (n = 2,208) in the context of IPTp clinical trials (20–22). Among them, 993 were from Mozambique, 854 from Benin, 131 from Gabon, 296 from Kenya, and 854 from Benin, 131 from Gabon, 296 from Kenya.

Table. Participants in study of VAR2CSA serologic testing to detect Plasmodium falciparum transmission patterns, by country and HIV status*

<table>
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<tr>
<td></td>
<td>Mozambique, n = 65</td>
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<td>Benin, n = 854</td>
<td>Gabon, n = 131</td>
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<td>188 (22)</td>
<td>38 (29)</td>
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<td>304 (63)</td>
<td>666 (78)</td>
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<tr>
<td>&lt;20</td>
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<td>281 (33)</td>
<td>45 (34)</td>
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<td>≥25</td>
<td>29 (45)</td>
<td>181 (37)</td>
<td>487 (57)</td>
<td>44 (34)</td>
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<td>Sulfadoxine/</td>
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<td>151 (31)</td>
<td>288 (34)</td>
<td>55 (42)</td>
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<td>Positive</td>
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<td>13 (3)</td>
<td>110 (15)</td>
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<tr>
<td>Negative</td>
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<td>468 (97)</td>
<td>616 (85)</td>
<td>125 (98)</td>
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<td>qPCR‡</td>
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<tr>
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<td>332 (46)</td>
<td>9 (10)</td>
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<td>49 (75)</td>
<td>424 (94)</td>
<td>393 (54)</td>
<td>80 (90)</td>
</tr>
</tbody>
</table>

*1PPT, intermittent preventive treatment during pregnancy; qPCR, quantitative PCR. 14% (196/1365) of HIV-uninfected and 12% (43/362) of HIV-infected participants were pregnant women with samples collected also at recruitment and second IPTp administration.
†All HIV-infected women who received placebo were also receiving cotrimoxazol prophylaxis.
§Maternal microscopic infection defined by the presence of P. falciparum parasites in peripheral blood or in placenta on microscopic or histologic examination, respectively.
¶Not determined: 178 microscopy and 196 qPCR.
*Maternal qPCR-positive infection was defined by a positive result on qPCR testing in peripheral or placental blood.
31 from Tanzania, and 49 from Spain. The baseline characteristics of the women selected for this trial were similar to those of the 6,216 women participating in the randomized clinical trials (Appendix 2 Table 1, https://wwwnc.cdc.gov/EID/article/25/10/18-1177-App2.xlsx).

The study areas represented 5 sites in sub-Saharan Africa with different intensities of malaria transmission. Prevalence of \textit{P. falciparum} infection detected by qPCR at delivery, in either peripheral or placental blood (averaged for 2010–2012), among HIV-uninfected women was 46% (332/725) in Benin, 10% (9/89) in Gabon, and 6% (28/452) in Mozambique and among HIV-infected women was 8% (22/273) in Kenya and 4% (13/327) in Mozambique (Table). The prevalence of \textit{P. falciparum} infection among pregnant women in Mozambique decreased from 25% (37/146) in 2004–2005 to 2% (3/176) in 2010 and increased to 6% (4/72) in 2012. A subset of 239 pregnant women from Mozambique recruited during 2011–2012 was followed during pregnancy; prevalence of \textit{P. falciparum} infection detected by qPCR was 16% (38/239) at first antenatal visit (mean gestational age ± SD, 20.7 ± 5.45 weeks), 3% (8/239) at the second IPTp administration (25.9 ± 4.98 weeks), and 5% (13/239) at delivery (38.4 ± 2.26 weeks).

\textit{P. falciparum} infection was detected at unscheduled visits for 2% (5/239) of the women. Overall, \textit{P. falciparum} infection was detected at any of these time points for 21% (49/239) of the women.

\textbf{\textit{P. falciparum}–Specific Antibody Profiles and Parasite Exposure during Pregnancy}

Mean antiparasite IgG levels in pregnant women from Mozambique delivering from 2010 through 2012 were above levels against bovine serum albumin plus 3 SD and higher than IgG levels in pregnant women from Spain except for DBL6ε and 3 of 25 VAR2CSA peptides (Figure 1, panel A; Appendix 2 Table 3). Five VAR2CSA peptides, DBL6ε, and pCSP were recognized by IgG from ≥5% of the pregnant women from Spain who had never been exposed to \textit{P. falciparum} (Figure 1, panel B), suggesting unspecific recognition; thus, these peptides were excluded from subsequent analysis. To further narrow down the VAR2CSA peptide candidates, we compared IgG levels in pregnant women from Mozambique delivering in 2004–2005 and 2010–2012, a period when \textit{P. falciparum} prevalence assessed by qPCR at delivery in peripheral or placental blood dropped from 25% to 5% (Figure 1, panel C) (3). This
decline in infection rates was mirrored by drops of IgG levels against 10 of the 18 previously selected VAR2CSA peptides (p1, p5, p8, p10, p12, p20, p27, p36, p38, p39) (Figure 1, panel D; Appendix 2 Table 4).

**Acquisition and Decay of IgG Responses against VAR2CSA**

We assessed the dynamics of IgG responses in a longitudinal cohort of 239 pregnant women from Mozambique (Figure 2, panel A). At delivery, compared with uninfected women, the 49 (21%) women infected with *P. falciparum* during pregnancy had higher IgG levels against the 10 down-selected peptides (Figure 2, panel B; Appendix 2 Table 5). At delivery, seroprevalence rates for p1 (23%), p5 (26%), p8 (26%), and p39 (31%) antibodies were above the cumulative prevalence of *P. falciparum* infection during pregnancy (Figure 2, panel C; Appendix 2 Table 5). No difference in IgG levels was observed between primigravid and multigravid women (Figure 2, panel D; Appendix 2 Table 5). The time to double (*T*₂ₓ) after *P. falciparum* infection ranged from 0.45 years (95% CI 0.31–0.80 years) for p5 to 1.07 years (95% CI 0.60–5.23 years) for p27 (Figure 2, panel E; Appendix 2 Table 5).

**Figure 2.** IgG responses during pregnancy against selected VAR2CSA antigens and polymorphism in target sequences in serologic study of *Plasmodium falciparum* in pregnant women. A) *P. falciparum* prevalence by quantitative PCR (qPCR) in 239 pregnant women from Mozambique at recruitment, second administration of IPTp, and delivery. Cumulative prevalence at delivery refers to peripheral or placental infection detected by microscopy, qPCR, or histology at any time point. B) Ratio of nMFIs at delivery in women from Mozambique infected during pregnancy compared with uninfected women. Error bars indicate 95% CIs. C) Seroprevalence at delivery, showing the cumulative prevalence of infection during pregnancy (red dashed line) and the prevalence at delivery by qPCR (light blue line). D) Ratio of nMFIs at delivery in multigravid compared with primigravid women, adjusted by IPTp, parity, age, and HIV status. Error bars indicate 95% CIs. E) IgG dynamics during pregnancy with estimates of time to double (*T*₂ₓ) and half-life (*T*₁/₂) obtained from linear mixed-effect regression model. Red points represent *P. falciparum* infection, dark gray lines the seropositivity cutoff, red lines the fitted-estimation, and dashed lines the 95% CI. F) Space-feeling representation of DBL1X-ID1 showing p5 (blue) and p8 (red). G) Logo representation of p5 and p8 sequences obtained from 50 *P. falciparum* isolates (20 from Mozambique, 10 from Benin, 10 from Gabon, and 10 from Kenya). IPTp, intermittent preventive treatment during pregnancy; nMFI, normalized median fluorescent intensity.
2 Table 6). IgG half-life among seropositive women at recruitment without evidence of *P. falciparum* infection during follow-up ranged from 0.55 (95% CI 0.38–1.02) years for p8 to 3.66 (95% CI 0.98–∞) years for p1 (Figure 2, panel E; Appendix 2 Table 6). Among recombinant antigens, IgG DBL5Ɛ showed the lowest T2 (0.31 [95% CI 0.21–0.61] years) and half-life (0.66 [95% CI 0.42–1.65] years), whereas AMA1 IgG showed the highest T2 (1.76 [95% CI 0.76–∞] years) and half-life (4.18 [95% CI 1.86–∞] years).

Among the down-selected VAR2CSA peptides (p1, p5, p8, and p39), IgG against p5 (51 amino acids) and p8 (48 amino acids) showed the lowest half-lives (0.55 [95% CI 0.38–1.02] years for p8; 1.33 [95% CI 0.65–∞] years for p5) and the largest increase in women exposed to *P. falciparum* during pregnancy compared with uninfected women (adjusted ratio [AR]p5 2.15 [95% CI 1.39–3.31] and ARp8 2.17 [95% CI 1.46–3.23]; Figure 2, panel B; Appendix 1 Figure 5; Appendix 2 Table 5). IgG levels and seroprevalence rates at delivery for p5 and p8 were higher among pregnant women with active or past malaria infection than among women with no parasite or pigment in the placenta, as assessed by histologic examination (Appendix 2 Table 7). 3D modeling mapped both sequences on the exposed surface of DBL1X-ID1 region of VAR2CSA (Figure 2, panel F). Amino acid variability obtained from 50 *P. falciparum* isolates collected at study sites was 5% ± 2 SD for p5 sequences and 16% ± 5 SD for p8 sequences, compared with the consensus peptide sequence included in the array (Figure 2, panel G; Appendix 1 Figures 3, 4).

**Performance of Selected VAR2CSA Peptides for Assessing Spatial and Temporal Differences in *P. falciparum* Exposure**

In pregnant women from Mozambique at delivery, p5 and p8 seroprevalence rates, as well as the composite of both (p5+8), decreased from 2004–2005 to 2010 (adjusted odds ratio [AOR]p5 0.27 [95% CI 0.11–0.68]), followed by an increase from 2010 to 2012 (AORp5 2.49 [95% CI 1.34–4.61]; Figure 3, panel A; Appendix 2 Table 8). This decrease and subsequent increase mirrored *P. falciparum* prevalence by qPCR. HIV infection and parity did not modify the associations observed (p value for interaction >0.05 for all cases; Appendix 2 Table 8). Similar to *P. falciparum* prevalence determined by qPCR, seroprevalence rates were the highest in HIV-uninfected women from Benin, followed by those from Gabon (AORp5 0.31 [95% CI 0.21–0.47]) and Mozambique (AORp5 0.21 [95% CI 0.16–0.28]; Figure 3, panel B; Appendix 2 Table 9). At delivery, pregnant women living in an area from Tanzania where no *P. falciparum* infection was detected by qPCR were seronegative against p5, p8, and p5+8 antibodies; 42% were seropositive against AMA1 and 48% were seropositive against MSP119 antibodies (Figure 3, panel B). Among HIV-infected women, seroprevalence rates for p8 and p5+8 were lower in Mozambique than in Kenya (AORp5 0.58 [95% CI 0.38–0.88]; Figure 3, panel C; Appendix 2 Table 9). p5 and p5+8 seroprevalence rates were higher among anemic than among nonanemic women (AORp5 1.26 [95% CI 1.03–1.55]; Figure 3, panel D; Appendix 2 Table 10). Seroprevalence rates were lower among HIV-uninfected women who received IPTp with mefloquine than among those who received sulfadoxine/pyrimethamine (AORp5 0.74 [95% CI 0.59–0.94]; Figure 3, panel E; Appendix 2 Table 11). Seroprevalence rates among HIV-infected women were lower among those who received mefloquine than among those who received placebo, although differences were not significant (AORp5 0.76 [95% CI 0.50–1.15]; Figure 3, panel F; Appendix 2 Table 11).

**Geographic Patterns of *P. falciparum* Transmission through VAR2CSA Serologic Testing**

Spatial geocoordinates were available for 698 pregnant women from Mozambique residing in Manhiça District (southern Mozambique). Geographic areas experiencing significantly higher seroprevalence rates than would be expected by chance were observed for p5 (radius 2.82 km; p = 0.024) and p5+8 (radius 1.06 km; p = 0.049) but not for MSP119 and AMA1 (Figure 4; Appendix 2 Table 12). The distribution of HIV infection, parity, age, and IPTp was similar among women inside and outside the serologic hotspot (p>0.05; Appendix 2 Table 12).

**Discussion**

Routine *P. falciparum* testing of easily accessible pregnant women at maternal healthcare services has the potential to offer a rapid, consistent, and cost-effective method for evaluating the malaria burden in different communities and tracking progress of interventions. IgGs against 2 VAR2CSA peptides, selected according to their ability to maximize the information about recent *P. falciparum* exposure during pregnancy, reflected differences in malaria burden over time and space in multiple settings in Africa and changes in parasite rates associated with the use of different preventive regimens. Overall, our results indicate that in areas with well-attended maternal healthcare services, this pregnancy-specific serologic test may serve as a useful sentinel surveillance tool for flagging changes in malaria burden and progress in the path toward elimination.

p5 (51 amino acids) is localized in the DBL1X domain and p8 (48 amino acids) in the ID1 region of VAR2CSA. Limited diversity (5%) of p5 sequence was observed in *P. falciparum* isolates from a variety of regions of Africa, in accordance with estimates from previous studies for the DBL1X domain (24). p8 corresponds to a more diverse (16%) variant of the ID1 region in VAR2CSA (25).
peptides are exposed on the DBL1X-ID1 N terminal region of VAR2CSA (18,26) and recognized by IgG from malaria-exposed pregnant women at levels higher than those of pregnant women from Spain and men from Mozambique (19). IgG responses against both VAR2CSA peptides increased with *P. falciparum* infection during pregnancy. Moreover, higher risk for anemia among p5 and p5+8 sero-responders support these antibodies as markers of recent infection, which adversely affects the women’s health (3). In contrast to the slow decay of IgG responses against AMA1, the half-life of IgG against p5 and p8 was <2 years, the average time reported in Mozambique for a second pregnancy to occur (27). The short half-life of p5 and p8 IgG, together with the similar IgG levels in multigravid and primigravid women, suggests that antibodies acquired during one pregnancy are not maintained over multiple pregnancies; thus, antibodies can be used as a reliable indicator of recent exposure for pregnant women, regardless of parity.

Seroprevalence rates for p5, p8, and the composite of both peptides (p5+8) mirrored trends in *P. falciparum* prevalence among pregnant women from Mozambique delivering during 2004–2012 (3), a temporal pattern that was also
observed for PfPR2–10 (28). Trends were similar among HIV-uninfected and infected women, suggesting that impairment of *P. falciparum*-specific antibody responses driven by viral infection (29) may not affect short-lived IgG responses against p5 and p8. Seroprevalence also reflected the burden of malaria among pregnant women residing in a variety of settings in Africa, as well as reductions in infection rates resulting from the use of mefloquine as IPTp among HIV-uninfected women (21). Similar trends, although not statistically significant, were observed among HIV-infected women receiving cotrimoxazole prophylaxis alone or in combination with mefloquine (22), possibly because of the longer duration of protection provided by 3 IPTp doses in HIV-infected women compared with the 2 doses in HIV-uninfected women. We also found that pregnant women living in an area from Tanzania where no *P. falciparum* infection was detected by qPCR as well as pregnant women from Spain never exposed to malaria were seronegative against p5 and p8, suggesting that pregnancy-specific serology might be used to confirm the eventual interruption of transmission.

Geographic distribution of pregnant women from Mozambique who were seropositive against p5 and p5+8 revealed a serologic hotspot in an area close to the river and sugar cane plantations, where the density of anopheline mosquitoes can be expected to be higher. In contrast, antibodies against MSP119 and AMA1 were not able to identify these malaria transmission patterns because of saturation of antibody responses after lifelong exposure to *P. falciparum*. These results support the value of using VAR2CSA serologic testing to amplify signals of recent exposure and suggest its potential to trigger targeted interventions to persons living in close proximity to passively detected seropositive pregnant women.

Our study has several limitations. First, the peptide array we used may have missed some conformational non-linear epitopes. Second, different transmission dynamics and host genetic factors may affect the acquisition and decay of antibodies (16). Third, steeper decay of antibodies may be observed out of pregnancy when infecting parasites express non-VAR2CSA variants. Fourth, the reduction of data from median fluorescence intensity to seroprevalence to simplify the serologic information of the assay may reduce the depth of serologic information. Developing alternative mathematical models that use antibody levels (30) may increase the sensitivity to detect temporal and spatial changes in malaria transmission. Last, antibody assessments in this study were conducted mainly at delivery; further studies should assess the performance of this testing at antenatal visits or soon after delivery (i.e., during infant immunization). Future research is needed to describe the relationship between pregnancy-specific serologic testing and malaria transmission in the general population and its value for confirming interruption of malaria transmission and providing early signals of *P. falciparum* resurgence after local elimination.

In summary, this study shows that IgG against 2 VAR2CSA peptides from the DBL1X-ID1 domain reveal...
temporal and spatial differences in malaria burden among pregnant women and reductions in exposure associated with the use of preventive measures during pregnancy. These antibodies enable the identification of local clusters of transmission that are missed by detection of \textit{P. falciparum} infections. Our results suggest that inferring recent exposure through VAR2CSA serologic testing would amplify signals of ongoing malaria transmission and increase the power to detect changes, either natural or driven by deliberate efforts, as well as malaria hotspots, among pregnant women (2). Moreover, peptides such as p1 targeted by long-lasting IgG responses may be useful for capturing past changes in transmission by sampling women of child-bearing age and relating seroprevalence with the number and timing of previous pregnancies. Operationally suitable serologic tests (31) capable of detecting antibodies against VAR2CSA synthetic peptides may be used in programmatic environments to stratify areas based on malaria burden, measure the effects of interventions, and document year-to-year changes in transmission.

Acknowledgments

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A patent application has been filed for the use of p5 and p8 for serologic surveillance (US 376 patent application no. 62523828, filed on June 23, 2017, by A.M.).

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References


VAR2CSA Serology to Detect *Plasmodium falciparum* Transmission Patterns

Appendix 1

**P. falciparum antigens**

A total of 46 peptides (length between 35 and 65 aminoacids) from conserved and semi-conserved regions of the VAR2CSA protein were designed after alignment by Clustal W of 18 VAR2CSA full-length sequences from *P. falciparum* isolates of different geographic origins (Asia, Africa, Central and South America) (*I*). The 25 peptides analyzed (3 peptides from DBL1X, 3 from DBL2X, 5 from DBL3X, 2 from DBL4Ɛ, 5 from DBL5Ɛ, 2 from DBL6Ɛ and 5 from NTS and ID regions) were previously selected based on highly recognized by IgGs from plasma of malaria exposed pregnant women compared with Spanish individuals and malaria exposed Mozambican men (*I*) (Appendix Table 2). A circumsporozoite peptide (pCSP) of 64 aminoacids (NVDP[NANP]_{15}) was also analyzed. Peptides were synthetized by Gl Biochem (Xangai, China) and median purity was estimated as 79% (range: 71%–91%) by high performance liquid chromatography (HPLC) and mass spectrometry. The Duffy binding-like recombinant domains (DBL3X, DBL5Ɛ and DBL6Ɛ) and the *P. falciparum* general antigens (AMA1 and MSP1_{19}) were all produced at ICGEB, New Delhi, India. *Clostridium tetani*, tetanus toxin was purchased from Santa Cruz Biotechnology (Dallas, Texas) to control for the amount of IgGs eluted from dried blood spots (DBS).

**Bead-based immunoassay**

Two multiplex suspension array panels were constructed to quantify IgG responses against *P. falciparum* recombinant proteins and synthetic peptides, using the xMAP technology and the Luminex® 100/200 System (Luminex® Corp., Austin, Texas). MagPlex® microspheres (magnetic carboxylated polystyrene microparticles, 5.6 μm) with different spectral signatures were selected for each protein (DBL3X, DBL5Ɛ, DBL6Ɛ, AMA1 and MSP1_{19}), peptide (25
VAR2CSA peptides and pCSP), tetanus toxin and bovine serum albumin. Antigens were covalently coupled to beads following a modification of the Luminex® Corporation protocol (1). Briefly, 200 μl of beads (2.5x10⁶) were transferred into a 1.5 mL eppendorf tube and resuspended by sonication and vortexing. The supernatant was removed after precipitation of the beads by magnetic separation during 60 seconds. Beads were washed twice with 250 μl of distilled water and pellets were resuspended in 80 μl of activation buffer (0.1 M NaH2PO4, pH 6.2). Sulfo-NHS (N-hydroxysulfosuccinimide) and EDC (1-Ethyl-3-[3-dimethylaminopropyl] carbodiimide hydrochloride; Pierce, Thermo Fisher Scientific Inc., Rockford, IL) were simultaneously added to reaction tubes at 5 mg/mL each in activation buffer, and reaction tubes were incubated at room temperature with gentle agitation, protected from light for 20 minutes. Activated beads were washed twice with 250 μl of coupling buffer (MES 50 mM, 2-[N-morpholino] ethanesulfonic acid monohydrate pH 5, Sigma-Aldrich). Antigen amounts in the coupling reaction for one million beads were determined after a titration experiment, and were as follows: 2 μg of tetanus toxin, 4 μg of DBL3X, DBL5Ɛ, DBL6Ɛ, AMA1, 8 μg of MSP119, 170 μg of each peptide and for bovine serum albumin a 1% solution in PBS (Phosphate-buffered saline) and volume was adjusted with coupling buffer until a maximum of 400 μl. Beads and antigens were vortexed, sonicated and then incubated over night at 4°C in the dark, with shaking. Coupled beads were blocked with 500 μl of 1% BSA in PBS for 30 minutes on a shaker at room temperature, avoiding light and then washed twice with 500 μl of assay buffer (1% BSA, 0.05% sodium azide in PBS filtrated) and resuspended in 400 μl of the same buffer. Beads were quantified on a Guava PCA desktop cytometer (Guava, Hayward, CA), and stored at 4°C in the dark. Protein and peptide multiplex arrays were prepared by pooling together equal volumes of coated beads.

Immediately before use, stock suspensions of antigen-coated microspheres were thoroughly resuspended by vortexing and sonication. Frozen plasma or the product of DBS elution were thawed at room temperature, mixed by vortexing, and spun at 16000 g for 5 minutes to remove particles. 50 μl of diluted microspheres (1000 microspheres/analyte/well) were added to a 96-well Mylar flat-bottom plate following the addition of 50 μl of diluted sample in duplicates to a final concentration of 1:400 for protein array and 1:100 for peptide array and incubated for 1 hour in agitation, protected from light at room temperature. After incubation, the plates were washed three times with 200 μl of washing buffer (0.05% Tween 20 in PBS) by
pelleting in a magnetic 96 well separator. 100 μl of biotinylated anti-human IgG (Sigma, Tres Cantos, Spain) diluted 1:1000 in assay buffer was added to each well, and plates were incubated for 45 minutes in agitation, protected from light at room temperature. After the incubation period, the plates were washed and 100 μl of streptavidin-conjugated R-phycoerythrin (Invitrogen, Carlsbad, CA) at a 1:1000 dilution in assay buffer was added and incubated for 25 minutes. Finally the plates were washed as before and the beads were resuspended in 100 μl of assay buffer and analyzed using the Luminex® 100/200 System.

A hyperimmune plasma pool composed by 23 plasmas from P. falciparum infected Mozambican pregnant women was tested to determine if the coupling was effective and was included in each assay plate as positive control, in addition to blanks (wells without sample) to assess background levels. A minimum of 50 microspheres were read per spectral signature and results were exported as crude median fluorescent intensity (MFI). Duplicates were averaged and background MFIs were subtracted. A total of 224 plates were analyzed and the intra-assay variation (mean CV of replicates from 20 plasma samples per plate) ranged from 1.4% to 7.3% for the protein array and from 2.5% to 12.4% for the peptide array. The inter-assay variation (variability of positive pool between 224 plates) was 5% for the protein array and 26% for the peptide array (1). To assure the validity of the luminex plates, a quality control was performed on the MFI values of the positive control pool. Results were plotted in Levey Jenning Charts (Appendix Figure 1) and five plates from the peptide array fell out of −2 standard deviation (SD) and +2SD and were re-analyzed (2). Results were normalized (nMFI) to account for plate-to-plate variation by dividing the background subtracted MFI of each sample by the value of the positive pool in the same plate and multiplying by the median of positive pools in all plates.

**Definition of IgG seropositivity**

Seropositivity cutoffs were obtained using finite mixture models (FMM) for pregnancy-specific malaria antigens (VAR2CSA peptides and recombinant domains). FMM can be applied in scenarios of heterogeneous IgG distributions where a subgroup of the study population is seronegative and another seropositive avoiding additional sampling of malaria never exposed pregnant women. However, seropositivity cutoffs to general malaria antigens (AMA1, MSP119 and pCSP) were obtained from means plus three SD of IgG levels from never exposed pregnant
women as a consequence of absence of heterogeneous IgG distribution against *P. falciparum* general antigens in malaria exposed adult pregnant women (1).

**Reconstitution of dried blood spots**

Antibodies were eluted from a total of 880 DBS from Gabon (n = 310), Kenya (n = 408) and Tanzania (n = 162) as previously described (1,3). Briefly, four spots of ≈3 mm in diameter were cut from the filter papers using a punch (McGill® round punch, 3 mm) and transferred to individual wells of a 96-well polystyrene U-bottom plate. Antibodies were eluted with 200 μl Luminex® assay buffer (1% BSA, 0.05% sodium azide in filtrated PBS) at room temperature overnight with gentle mixing which, assuming a hematocrit of 50%, gives a concentration of eluted blood proteins equivalent to a 1:50 plasma dilution (3,4).

To assess the quality of the elution, hemoglobin levels in the eluted DBS were measured by spectrophotometry (wavelengths 415, 380 and 450) and calculated using the Harboe method with the Allen correction (Hb [mg/l] = 167.2 x A415 – 83.6 x A380 – 83.6 x A450) x dilution factor). Three criteria to discard DBS improperly eluted were followed, as previously described (1). First, 259 reddish-brown spots against a pale background were discarded after visual examination of reconstituted spots.(3) Second, 10 DBS were also discarded because hemoglobin levels measured in the elutions were below the upper quartile of hemoglobin value among samples considered with inappropriate visual aspect (hemoglobin upper quartile = 7.4m/l). Finally, 153 samples were also discarded because anti-tetanus antibodies measured in the elutions were below the lowest quartile obtained from anti-tetanus IgG among samples with appropriate visual aspect and hemoglobin levels (anti-tetanus lowest quartile = [11563,5 nMFI]). Finally, 458/880 samples (131 from Gabon, 296 from Kenya and 31 from Tanzania) were considered as correctly eluted.

**Sequencing of var2csa p5 and p8 in *P. falciparum* isolates**

A total of 50 *P. falciparum* isolates collected on filter paper (Whatman 903) from infected individuals in Mozambique (n = 20), Benin (n = 10), Gabon (n = 10) and Kenya (n = 10) were selected for DNA sequencing. A half of the filter paper containing a 25 μL of blood drop was used for DNA extraction using a QIAamp DNA Mini kit (Qiagen), as per the manufacturer's
instructions. Finally, DNA was eluted in 100 μL of AE buffer given in the kit. The presence of *P. falciparum* infection was detected using a previously described method (5,6).

We designed a single polymerase chain reaction (PCR) based assay to amplify purified DNA templates using 2720 Thermal Cycler (Applied Biosystems) followed by Sanger sequencing for *var2csa* gene. In brief, a 25 μl reaction was set up, containing 0.5 μM of each forward (p5_F- 5’aaggttggaagttattac-3’) and reverse (p8_R- 5’attagtaagatgcaagtact-3’) primers, 1x HOT FirePol Master Mix (Solis BioDyne; Cat. No. 04–27–00125) and 5 μl of template DNA. The reaction volume was raised by PCR-grade water. The template DNA was denatured at 95°C for 15 minutes in a thermocycler, followed by 35 cycles of amplification (95°C for 1 minute, 54°C for 1 minute, and 72°C for 1 minute) and a final extension at 72°C for 10 minutes. A reaction using 5 μl of PCR-grade water instead of template DNA was included as a negative control. PCR products were run on 1.5% agarose (Invitrogen) gels in 1× TBE buffer (Tris/Borate/EDTA; Thermo Fisher Scientific) to determine the presence and size of the amplified DNA. PCR products were visualized using a ultraviolet trans-illuminator. The PCR primer set was also tested with human gDNA to check their specificity. The expected size of the PCR was 960 bp covering amino acid positions 220 to 539 of *var2csa* gene. PCR products were quantified using EPOCH Biotech system. Approximately 1200 ng of PCR products were sent to Genewiz, following safety instructions for the accurate shipment of PCR amplicons. To sequence the p5 and p8 fragments of *var2csa* gene, three new sequencing primers were used (Seq_p5_R - 5’ccatttcttcacacattcac-3’; Seq_p8_F - 5’gggtgatctttcgcagaa-3’; Seq_p8_R-5’cgcaagaatctttggaacaca-3’) along with PCR p5_F primer. This allowed us to sequence 960 bp *var2csa* amplicons in both directions. The bi-directional sequencing with PCR p5_F and Seq_p5_R, and Seq_p8_F and Seq_p8_R primer sets covered 220–335 and 389–499 aa respectively.

The variations in the test sequences were identified by sequence alignment (Blastn, NCBI:https://blast.ncbi.nlm.nih.gov) against reference sequence of 3D7 (PF3D7_1200600) retrieved from PlasmoDB. The nucleotide sequences obtained from field isolates were translated using ExPASy online tool (http://web.expasy.org/translate/) and represented as a logo figure using the Weblogo Version 2.8.2 online tool (http://weblogo.berkeley.edu).
VAR2CSA DBL1X-ID1 3D model

The 3D-structure of DBL1X-ID1 was obtained by submitting the 3D7 sequence, with domain limits defined by Bockhorst and colleagues (2007) (7) to the HHpred server (http://toolkit.tuebingen.mpg.de/hhpred). The structure with highest HHpred score, corresponding to the DBL1α domain of the VarO strain (Protein Data Bank [PDB] 2yk0 (8)), was selected for homology modeling in MODELER based on the default alignment. Molecular graphics were generated in UCSF Chimera version 1.5.3 (9).

IgG dynamic analysis through mixed-effects regression models

Summary statistics according to the longitudinal design were calculated by trimester. Time-at-risk was estimated using gestational age at recruitment as the time when subjects first came under observation, and gestational age at delivery as the latest time under which the subjects were both under observation and at risk. Antibody levels were analyzed assuming a lognormal distribution, and therefore they were described by the geometric means and the overall, between and within-subjects standard deviations.

The crude and adjusted effect of *P. falciparum* infection on antibody levels was analyzed using log-linear mixed-effects regression models incorporating Gaussian random intercepts. This resulted in an estimate of the rates of antibody dynamics (boosting or decay), assuming a single exponential model. Time to 2-fold increase were calculated from the estimated rates and the boundaries at 95% confidence interval (CI) obtained from mixed-effects models for women with *P. falciparum* infection at follow-up (10,11). Similarly, half-lives were calculated from models including women who were seropositive at recruitment with no *P. falciparum* infection at follow-up (10,11). Where the boosting rate is a negative value (rate below 1) or the decay rate is a positive value (rate above 1), the calculated time to 2-fold increase or half-life was reported as infinity. Statistical comparisons were performed at two-sided significance level of 0.05 and 95% CI were calculated for all estimations.

Consider our longitudinal dataset consisting of antibody measurements of 239 pregnant women on three successive gestational ages. Because we were not really interested in these particular 239 women per se, we treated them as a random sample from a larger population and
modeled the between-woman variability as a random effect, as a random-intercept term at the woman level. We thus fitted the model:

\[
\ln(C_{ij}) = \beta_0 + \beta_1 T_{ij} + u_j + e_{ij}
\]

where \(C_{ij}\) are the concentrations for \(i = 1 \ldots\) Three measurement of gestational age (T) and \(j = 1 \ldots 239\) women. The fixed portion of the model, \(\beta_0 + \beta_1 T_{ij}\), simply states that we wanted one overall regression line representing the population average. The random effect \(u_j\) serves to shift this regression line up or down according to each woman. Back-transforming the measurements to the original scale we obtain the following overall regression line:

\[
C_{ij} = e^{\beta_0 + \beta_1 T_{ij}} = e^{\beta_0} e^{\beta_1 T_{ij}} = C_0 e^{\beta_1 T_{ij}}
\]

where \(C_0\) is the baseline concentration. We can estimate the time required to obtain a value \(\Delta\) times \(C_{ij}\)

\[
\Delta C_{ij} = C_0 e^{\beta_1 (T_{ij} + \Delta)} = C_0 e^{\beta_1 T_{ij}} e^{\beta_1 \Delta} = C_{ij} e^{\beta_1 \Delta}
\]

and therefore

\[
\Delta = e^{\beta_1 \Delta}
\]

\[
\ln(\Delta) = \beta_1 \Delta
\]

\[
t_\Delta = \ln(\Delta) / \beta_1
\]

In particular, for half-life \(\Delta = 1/2\) and then \(t_{1/2} = \ln(1/2) / \beta_1 = -\ln(2) / \beta_1\). Similarly, time to 2-fold increase can be calculated as \(t_{2x} = \ln(2) / \beta_1\).

**Geospatial analysis**

Spatial hotspots of *P. falciparum* infection and seropositivity among pregnant women from Mozambique living in Manhiça district were designated using hierarchical cluster analysis with Ward’s minimum variance method (12). Kulldorff spatial scan method (13–15) was used to identify the most likely hotspot for *P. falciparum* infection and seropositivity, setting our significance threshold (\(\alpha\)) as 0.05. Hotspots of greater size than one cluster were identified by consecutively aggregating nearest-neighboring areas until a proportion of the total study population was included. To ascertain statistical significance, we employed Monte Carlo sampling, using Poisson likelihoods, and following the original Kulldorff method for
identification of clusters (16). Analysis were performed using the R statistical software (version 3.2.1) (17) and maps were generated using OpenStreetMap (18). The key R packages used were SpatialEpi (19), deldir (20), geosphere (21), rgeos (22) raster (23) and leaflet (24).

References


Appendix Figure 1. Levey Jenning Charts plotted for the quality control of the Luminex assay. Data analysis for a quality control of the MFI values obtained from the hyperimmune plasma pool (positive pool) for the A) protein array and B) peptide array. The dots represent each positive pool per plate. If these dots fell out of the mean of positive pool +/- two standard deviation (SD) (red area), these plates were rejected and re-analyzed.
*40% (196/485) of HIV-uninfected and 12% (43/362) of HIV-infected women participated in a longitudinal study in which samples were collected at recruitment, second IPTp administration and delivery (data available for qPCR, microscopy and histology).
Appendix Figure 3. VAR2CSA peptide p5 sequences from *P. falciparum* isolates from Mozambique, Benin, Kenya and Gabon. Grey fill means conserved regions, red indicate aminoacid change, blue indicate nucleotide change but same aminoacid and positions with two aminoacids correspond to mixed infections. Samples Mç1–10, B1–10, K1–10 and G1 were collected from pregnant women in Mozambique (Mç), Benin (B), Kenya (K) and Gabon (G) who were participating in the clinical trial (2010–2012). Samples MTP1–10 we obtained from Northern Mozambique (Montepuez).
Appendix Figure 4. VAR2CSA peptide p8 sequences from *P. falciparum* isolates from Mozambique, Benin, Kenya and Gabon. Grey fill means conserved regions, red indicate aminoacid change and blue indicate nucleotide change but same aminoacid. Samples Mç1–10, B1–10, K1–10 and G1 were collected from pregnant women in Mozambique (Mç), Benin (B), Kenya (K) and Gabon (G) who were participating in the clinical trial (2010–2012). Samples MTP1–10 we obtained from Northern Mozambique (Montepuez).
Appendix Figure 5. Diagram of peptide selection.