The Malaria TaqMan Array Card Includes 87 Assays for *Plasmodium falciparum* Drug Resistance, Identification of Species, and Genotyping in a Single Reaction

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**Abstract**

Antimalarial drug resistance exacerbates the global disease burden and compounds eradication efforts. To facilitate the surveillance of resistance markers in countries of malaria endemicity, we developed a suite of TaqMan assays for known resistance markers and compartmentalized them into a single array card (TaqMan array card, TAC). We included 87 assays for species identification, for the detection of *Plasmodium falciparum* mutations associated with chloroquine, atovaquone, pyrimethamine, sulfadoxine, and artemisinin resistance, and for neutral single nucleotide polymorphism (SNP) genotyping. Assay performance was first optimized using DNA from common laboratory parasite lines and plasmid controls. The limit of detection was 0.1 to 10 pg of DNA and yielded 100% accuracy compared to sequencing. The tool was then evaluated on 87 clinical blood samples from around the world, and the malaria TAC once again achieved 100% accuracy compared to sequencing and in addition detected the presence of mixed infections in clinical samples. With its streamlined protocol and high accuracy, this malaria TAC should be a useful tool for large-scale antimalarial resistance surveillance.

**Keywords** TaqMan PCR, antimalarial resistance, malaria, mutation, surveillance tool

Despite a decline in incidence, malaria remains an important cause of morbidity and mortality across the world. There were 214 million cases of malaria globally, leading to an estimated 429,000 to 730,500 deaths in 2015 (1, 2). Antimalarial drug resistance has substantial implications for malaria control. The emergence of resistance to chloroquine in the 1980s led to its replacement with sulfadoxine-pyrimethamine (SP) for the treatment of uncomplicated malaria (3). The rapid development of resistance to these drugs then led to the recommendation of artemisinin combination therapy (4). Recently, resistance to artemisinin has been detected in the Greater Mekong subregion including the China-Myanmar border (5–10). Because of this history, monitoring antimalarial resistance is an important component of successful malaria control programs.

Standard methodologies to assess antimalarial sensitivity are time-consuming, technically challenging, and expensive. To estimate clinical efficacy (*in vivo* sensitivity), the response to antimalarial treatment needs to be monitored in the patient for at least 14
days (11). Ex vivo assessment of Plasmodium falciparum sensitivity requires tissue culture facilities and fresh blood products for parasite propagation, which are not readily available in countries of malaria endemicity. Genetic markers of resistance have been identified for several clinical antimalarials, and assessment of parasite DNA for these markers is a useful method for resistance detection (12–20). Several molecular tools have been developed to detect these markers, including PCR-restriction fragment length polymorphism (RFLP) (21), high-resolution melt analysis (22), quantitative PCR (23, 24), and loop-mediated isothermal amplification (LAMP) (25). Although accurate, these PCR-based methods are onerous because multiple markers need to be assessed. Luminex-based assays (26, 27), microarrays (28), and whole-genome sequencing of clinical samples (29, 30) can achieve greater throughput but are difficult to implement in field settings. We therefore sought to create a comprehensive, easy-to-perform method to track many resistance markers from multiple samples in a single run. The TaqMan array card (TAC) is a customizable 384-well card that compartmentalizes each sample into 48 different quantitative PCRs. TAC protocols are streamlined (due to several self-contained reagents), and eight patient samples can be run simultaneously. Our group has applied this technology successfully for detection of multiple tuberculosis (TB) drug resistance markers (31, 32), syndromic pathogen detection (33, 34), and pneumococcal serotyping (35) and has found excellent reproducibility between multiple laboratories across Africa, Asia, and America.

Here, we describe the construction and testing of the malaria TAC. This initial card design includes 70 assays for the detection of resistance-associated mutations in the P. falciparum CRT (pfCRT chloroquine), pfCYTB (atovaquone), pfDHFR (pyrimethamine), pfDHPS (sulfadoxine), pfMDR1 (multidrug resistant), and pfKelch13 (artemisinin) genes. Additionally, assays for the detection of the five Plasmodium species targeting humans (falciparum, vivax, ovale, malariae, and knowlesi) and 12 genotyping single nucleotide polymorphisms (SNPs) were included to provide additional epidemiologic information about the samples.

RESULTS

TaqMan assay performance. The 35 duplex TaqMan assays for resistance markers, 6 duplex assays for SNP genotyping, and 5 singleplex assays for species identification as well as a control assay for human glyceraldehyde-3-phosphate dehydrogenase (hGAPDH) (Fig. 1) were first tested against 35 other blood pathogens, and no cross-reactivity (detection of false positives) was observed (see Materials and Methods for a list of pathogen) (data not shown). Specificity was then tested against 11 P. falciparum lines, P. vivax, P. knowlesi, human DNA, and 17 synthetic plasmid controls in the 384-well plate format (see Fig. S1 in the supplemental material). For most assays (38/47), we detected no cross-reactivity between any of the duplexed assays (wild-type and mutant loci and major and minor SNP alleles) or among the five Plasmodium species. We observed minor cross-reactivity in 9 of 47 assays (pfCRT 73-76VMET, pfCRT 326S, pfCRT 356I, pfCYTB 268Y, pfCYTB 268C, pfDHPS 613A, pfMDR1 1246Y, SNP1-T, and SNP2-G) (Fig. S1). Such cases were observed only when high concentrations of DNA (>1 ng) were used and were eliminated by adjusting the threshold.

Next, the PCR performance of each primer/probe assay was determined on 384-well plates and then later with the TAC format. To test the amplification efficiency of each assay, DNA from either individual parasite lines or plasmid controls was tested in triplicate for six serial dilutions (see Materials and Methods for details). The overall linearity ($R^2$) of the 88 targets, including hGAPDH, was 0.990 ± 0.01 and 0.994 ± 0.01; PCR efficiencies were 90% ± 6.4% and 92% ± 7.9% for the 384-well plates and TAC formats, respectively (Tables S1 and S2). The limit of detection was similar on both 384-well plates and TaqMan array cards, ranging between 10 to 100 plasmid copies per reaction or 4.03 to 403 copy equivalent genomic DNAs (0.1 to 10 pg) per reaction (Tables S1 and S2).

Malaria TAC evaluation. The performance of the TaqMan array card was evaluated using genomic DNA from 18 laboratory parasite lines and 87 clinical samples. The
malaria TAC successfully detected the correct wild-type or mutant loci in 99% ± 4.1% of reaction products using 18 laboratory parasite strains; those that were left undetermined represented parasite lines with mutations other than those included in malaria TAC (Table 1). The malaria TAC successfully detected either wild-type or mutant loci for 89% ± 7.3% of reactions using 87 clinical samples (Table 1). The pfCRT 72-76, pfCRT 97, pfDHFR 164, and pfDHPS 540 targets had the lowest rates of detection in this set of samples (≤80%) (Table 1). The undetermined samples were mostly from the Malawi repository, with an average detection level of 78% ± 9.0%. Thailand-, China-, and Uganda-derived samples exhibited detection levels of 95% ± 5.3%, 90% ± 18.4%, and 100%, respectively (Table S3). The malaria TAC successfully detected two clinical samples that contained non-<i>P. falciparum</i> parasite DNA, including one each of <i>P. vivax</i> and <i>P. ovale</i> (Table 1).

The allelic distribution of the loci successfully detected by the malaria TAC is shown in Fig. S2. The majority of clinical samples showed pfCRT mutations, except for those from Malawi. All samples displayed pfDHFR and pfDHPS mutations, but none were observed in pfCYTB. The number and type of pfMDR1 and pfKelch13 mutations varied depending on the country of origin. The pattern of <i>P. falciparum</i> mutant alleles detected in our analysis is summarized in Table S4. Overall, we detected 13 distinct resistance marker genotypes from 16 laboratory parasite lines (data not shown). This
### TABLE 1 Comparison of malaria TAC performance using laboratory parasites and clinical samples

| Target          | TAC performance on culture parasite lines (n = 18) | TAC performance on clinical samples (n = 87) |
|-----------------|--------------------------------------------------|-----------------------------------------------|
|                 | No. (%) of positive parasite lines (n = 18) | Mean C<sub>t</sub> ± SD | No. (%) of positive clinical samples (n = 87) | Mean C<sub>t</sub> ± SD |
| Human GAPDH     |                                                   |                                |                                                   |                                |
| P. vivax        | 1 (100)                                         | 24.2                           | 80 (92)                                         | 30.7 ± 3.4                    |
| P. knowlesi     | 1 (100)                                         | 19.4                           | 1 (100)                                         | 29.2                           |
| P. ovale        |                                                   |                                | 1 (100)                                         | 29.5                           |
| P. falciparum   | 16 (100)                                        | 21.9 ± 3.0                     | 81 (95)                                         | 29.4 ± 5.4                    |
| pfCRT 72–76     | 16 (100)                                        | 24.4 ± 2.6                     | 62 (73)                                         | 34.7 ± 4.3                    |
| pfCRT 97        | 15 (94)<sup>a</sup>                             | 25.9 ± 2.7                     | 56 (66)                                         | 37.6 ± 4.6                    |
| pfCRT 326       | 16 (100)                                        | 23.3 ± 2.9                     | 80 (94)                                         | 32.0 ± 4.9                    |
| pfCRT 356       | 16 (100)                                        | 24.0 ± 3.3                     | 80 (94)                                         | 31.7 ± 5.1                    |
| pfCYTB 258      | 16 (100)                                        | 18.5 ± 2.4                     | 85 (100)                                        | 25.2 ± 4.6                    |
| pfCYTB 268      | 16 (100)                                        | 18.1 ± 2.5                     | 81 (95)                                         | 26.6 ± 4.1                    |
| pfCYTB 272      | 16 (100)                                        | 19.9 ± 2.4                     | 83 (98)                                         | 27.6 ± 4.5                    |
| pfDHFR 51       | 16 (100)                                        | 25.9 ± 2.8                     | 72 (85)                                         | 35.7 ± 4.1                    |
| pfDHFR 59       | 16 (100)                                        | 23.8 ± 2.5                     | 76 (89)                                         | 33.4 ± 4.8                    |
| pfDHFR 108      | 16 (100)                                        | 25.6 ± 3.1                     | 70 (82)                                         | 36.0 ± 4.2                    |
| pfDHFR 164      | 16 (100)                                        | 23.2 ± 2.5                     | 66 (78)                                         | 33.9 ± 4.6                    |
| pfDHS436–437    | 12 (75)<sup>b</sup>                             | 24.1 ± 3.4                     | 79 (93)                                         | 30.8 ± 4.5                    |
| pfDHS540        | 16 (100)                                        | 25.8 ± 3.5                     | 61 (72)                                         | 36.1 ± 4.4                    |
| pfDHS581        | 16 (100)                                        | 24.5 ± 3.3                     | 73 (86)                                         | 32.5 ± 5.2                    |
| pfDHS613        | 16 (100)                                        | 23.2 ± 3.1                     | 77 (91)                                         | 30.2 ± 4.8                    |
| pfMDR1 86       | 16 (100)                                        | 23.9 ± 3.2                     | 78 (92)                                         | 32.2 ± 4.3                    |
| pfMDR1 184      | 16 (100)                                        | 24.2 ± 3.1                     | 74 (87)                                         | 33.3 ± 4.4                    |
| pfMDR1 1034     | 16 (100)                                        | 22.4 ± 3.2                     | 74 (87)                                         | 32.7 ± 5.1                    |
| pfMDR1 1042     | 16 (100)                                        | 22.5 ± 3.0                     | 74 (87)                                         | 33.4 ± 4.4                    |
| pfMDR1 1246     | 16 (100)                                        | 24.8 ± 3.1                     | 75 (88)                                         | 34.5 ± 4.7                    |
| pfKelch13 446   | 16 (100)                                        | 22.2 ± 2.9                     | 76 (89)                                         | 31.3 ± 4.8                    |
| pfKelch13 458   | 16 (100)                                        | 25.0 ± 2.7                     | 77 (91)                                         | 34.3 ± 4.6                    |
| pfKelch13 493   | 16 (100)                                        | 23.8 ± 3.0                     | 78 (92)                                         | 31.8 ± 4.6                    |
| pfKelch13 539   | 16 (100)                                        | 24.7 ± 2.9                     | 77 (91)                                         | 33.5 ± 4.3                    |
| pfKelch13 543   | 16 (100)                                        | 21.8 ± 2.8                     | 74 (87)                                         | 29.4 ± 4.6                    |
| pfKelch13 561   | 16 (100)                                        | 22.2 ± 2.8                     | 79 (93)                                         | 29.6 ± 4.6                    |
| pfKelch13 574   | 16 (100)                                        | 22.4 ± 2.8                     | 78 (92)                                         | 29.7 ± 4.6                    |
| pfKelch13 578   | 16 (100)                                        | 23.3 ± 2.8                     | 76 (89)                                         | 30.3 ± 4.7                    |
| pfKelch13 580   | 16 (100)                                        | 24.5 ± 3.1                     | 78 (92)                                         | 31.7 ± 5.1                    |
| Pf<sub>01</sub> 000130573 | 16 (100)                                      | 25.0 ± 2.8                     | 81 (95)                                         | 31.6 ± 5.0                    |
| Pf<sub>01</sub> 000539044 | 16 (100)                                      | 26.3 ± 3.0                     | 73 (86)                                         | 34.1 ± 5.4                    |
| Pf<sub>02</sub> 000842803 | 16 (100)                                      | 25.5 ± 3.1                     | 74 (87)                                         | 32.0 ± 5.3                    |
| Pf<sub>06</sub> 000145472 | 16 (100)                                      | 23.3 ± 2.9                     | 76 (89)                                         | 30.2 ± 4.7                    |
| Pf<sub>06</sub> 000937750 | 16 (100)                                      | 23.9 ± 3.2                     | 76 (89)                                         | 32.9 ± 4.4                    |
| Pf<sub>07</sub> 000277104 | 16 (100)                                      | 24.5 ± 3.0                     | 75 (88)                                         | 32.2 ± 4.5                    |

Mean % positive (±SD): (99.2 ± 4.1) for culture parasite lines and (89 ± 7.3) for clinical samples.

<sup>a</sup>C<sub>t</sub>, cycle threshold.

<sup>b</sup>One line was negative with both the wild-type probe (H) and mutant probe (Q) by TAC; sequencing mutant (L).

<sup>c</sup>Four lines were negative with both the wild-type probe (SG) and mutant probe (AG or SA); sequencing mutant (FG).

Estimation was more difficult for clinical samples since some targets were not detectable and therefore yielded incomplete data. However, from clinical samples in which all drug resistance loci were detected, the malaria TAC successfully detected 13 distinct resistance marker genotypes or antibiograms (data not shown).

For validation, Sanger sequencing was performed in parallel on all laboratory parasites and ~50% of clinical parasites. Results from the malaria TAC showed 100% sensitivity, specificity, and accuracy (Table 2). One laboratory parasite sample was negative for both wild-type (H) and mutant (Q) probes by malaria TAC at pfCRT 97. Sequencing revealed that this sample harbored an alternative allele that was not tested in our current design (H97L) (Tables 1 and 2). Similarly, four pfDHS436–437 mutants had an alternative allele (FG) that was not covered by the malaria TAC and was therefore not detected with the wild-type (SG) or mutant (AG or SA) probe (Tables 1 and 2).
TABLE 2 Comparison of malaria TAC and Sanger sequencing results

| Target    | TAC probe | Sequencing result | TAC sensitivity (%) | TAC specificity (%) | TAC accuracy (%) |
|-----------|-----------|-------------------|--------------------|---------------------|------------------|
|           |           | Mutant            | Wild type          |                     |                  |
| pfCRT 72–76 | Mutant    | 42                | 0                  | 100                 | 100              |
|           | Wild type | 0                 | 16                 |                     |                  |
| pfCRT 97  | Mutant    | 1                 | 0                  | 100                 | 100              |
|           | Wild type | 0                 | 57                 |                     |                  |
| pfCRT 326 | Mutant    | 36                | 0                  | 100                 | 100              |
|           | Wild type | 0                 | 22                 |                     |                  |
| pfCRT 356 | Mutant    | 35                | 0                  | 100                 | 100              |
|           | Wild type | 0                 | 23                 |                     |                  |
| pfCYTB 258| Mutant    | 0                 | 0                  | N/A                 | 100              |
|           | Wild type | 0                 | 58                 |                     |                  |
| pfCYTB 268| Mutant    | 1                 | 0                  | 100                 | 100              |
|           | Wild type | 0                 | 57                 |                     |                  |
| pfCYTB 272| Mutant    | 0                 | 0                  | N/A                 | 100              |
|           | Wild type | 0                 | 58                 |                     |                  |
| pfDHFR 51 | Mutant    | 49                | 0                  | 100                 | 100              |
|           | Wild type | 0                 | 9                  |                     |                  |
| pfDHFR 59 | Mutant    | 53                | 0                  | 100                 | 100              |
|           | Wild type | 0                 | 5                  |                     |                  |
| pfDHFR 108| Mutant    | 56                | 0                  | 100                 | N/A              |
|           | Wild type | 0                 | 2                  |                     |                  |
| pfDHFR 164| Mutant    | 27                | 0                  | 100                 | 100              |
|           | Wild type | 0                 | 31                 |                     |                  |
| pfDHP5 436/37 | Mutant | 13               | 0                  | 100                 | 100              |
|           | Wild type | 0                 | 45                 |                     |                  |
| pfDHP5 540 | Mutant    | 48                | 0                  | 100                 | N/A              |
|           | Wild type | 0                 | 10                 |                     |                  |
| pfDHP5 581 | Mutant    | 30                | 0                  | 100                 | 100              |
|           | Wild type | 0                 | 28                 |                     |                  |
| pfDHP5 613 | Mutant    | 4                 | 0                  | 100                 | 100              |
|           | Wild type | 0                 | 54                 |                     |                  |
| pfMDR1 86 | Mutant    | 10                | 0                  | 100                 | 100              |
|           | Wild type | 0                 | 48                 |                     |                  |
| pfMDR1 184 | Mutant    | 25                | 0                  | 100                 | 100              |
|           | Wild type | 0                 | 33                 |                     |                  |
| pfMDR1 1034 | Mutant | 2                 | 0                  | 100                 | 100              |
|           | Wild type | 0                 | 56                 |                     |                  |
| pfMDR1 1042 | Mutant | 5                 | 0                  | 100                 | 100              |
|           | Wild type | 0                 | 53                 |                     |                  |
| pfMDR1 1246 | Mutant | 6                 | 0                  | 100                 | 100              |
|           | Wild type | 0                 | 52                 |                     |                  |
| pfKelch13 446 | Mutant | 15                | 0                  | 100                 | 100              |
|           | Wild type | 0                 | 43                 |                     |                  |
| pfKelch13 458 | Mutant | 0                 | 0                  | N/A                 | 100              |
|           | Wild type | 0                 | 58                 |                     |                  |
During the assessment of clinical parasite genotypes, we found that 7/85 (8%) clinical DNA samples were positive for multiple probes at a single locus (termed hetero-resistance [data not shown]). These findings were originally detected using the malaria TAC but were subsequently sequence confirmed (Fig. 2 and S3). For example, DB133 was positive for both the wild-type (184Y) and mutant (184F) \textit{pfMDR1} probes; Sanger sequencing showed mixed T/A as position 184, which indicated the presence of both F(TTT) and Y(TAT) (Fig. 2).

Due to space limitations, only 6 of 24 original genotyping SNPs (36) could be included in this initial malaria TAC design. With this limited set, we detected 12 distinct barcodes from 16 laboratory parasite lines (data not shown) and 67 distinct barcodes from 85 clinical samples (Table 3) (18 samples exhibited incomplete data and were not included in this analysis). The set of distinct barcodes from clinical samples included 13 shared and 14 unique barcodes. Four of the 13 shared barcodes were found in multiple samples from the same country, 8/13 were shared across two different countries, and 1/13 (CATCAG) was found in all four countries. We also detected 14 (21%) barcodes that were unique and contained mixed alleles, which were also confirmed by sequencing (Fig. 2 and S4). In one example, sample DB009 was positive for both the major and

| Target       | TAC probe | Sequencing result | Mutant | Wild type | TAC sensitivity (%) | TAC specificity (%) | TAC accuracy (%) |
|--------------|-----------|-------------------|--------|-----------|--------------------|-------------------|-----------------|
| PfKelch13 493| Mutant    | 2                 | 0      | 100       | 100                | 100               | 100             |
|              | Wild type | 0                 | 56     |           |                    |                   |                 |
| PfKelch13 539| Mutant    | 1                 | 0      | 100       | 100                | 100               | 100             |
|              | Wild type | 0                 | 57     |           |                    |                   |                 |
| PfKelch13 543| Mutant    | 1                 | 0      | 100       | 100                | 100               | 100             |
|              | Wild type | 0                 | 57     |           |                    |                   |                 |
| PfKelch13 561| Mutant    | 1                 | 0      | 100       | 100                | 100               | 100             |
|              | Wild type | 0                 | 57     |           |                    |                   |                 |
| PfKelch13 574| Mutant    | 3                 | 0      | 100       | 100                | 100               | 100             |
|              | Wild type | 0                 | 55     |           |                    |                   |                 |
| PfKelch13 578| Mutant    | 0                 | 0      | N/A       | 100                | 100               | 100             |
|              | Wild type | 0                 | 58     |           |                    |                   |                 |
| PfKelch13 580| Mutant    | 2                 | 0      | 100       | 100                | 100               | 100             |
|              | Wild type | 0                 | 56     |           |                    |                   |                 |

\[\text{Minor allele} \quad \text{Major allele}\]

| Pf_01_000130573 | Minor allele (T) | 25 | 0 | 100 | 100 | 100 |
| Pf_01_000130573 | Major allele (C) | 0  | 33|     |     |     |
| Pf_01_000539044 | Minor allele (G) | 12 | 0 | 100 | 100 | 100 |
| Pf_01_000539044 | Major allele (A) | 0  | 46|     |     |     |
| Pf_02_000842803 | Minor allele (C) | 33 | 0 | 100 | 100 | 100 |
| Pf_02_000842803 | Major allele (T) | 0  | 25|     |     |     |
| Pf_06_000145472 | Minor allele (G) | 21 | 0 | 100 | 100 | 100 |
| Pf_06_000145472 | Major allele (C) | 0  | 37|     |     |     |
| Pf_06_000937750 | Minor allele (G) | 28 | 0 | 100 | 100 | 100 |
| Pf_06_000937750 | Major allele (A) | 0  | 30|     |     |     |
| Pf_07_000277104 | Minor allele (G) | 34 | 0 | 100 | 100 | 100 |
| Pf_07_000277104 | Major allele (A) | 0  | 24|     |     |     |

\(a\) Data represent both laboratory parasites (\(n = 16\)) and clinical samples (\(n = 42\)).

\(b\) Negative with both the wild-type probe (H) and mutant probe (Q) by TAC; sequencing mutant (L).

\(c\) Four of 13 samples were negative with both the wild-type probe (SG) and mutant probe (AG or SA); sequencing mutant (FG).

\(d\) N/A, not applicable.
DISCUSSION

In this study, we developed and tested a quantitative PCR-based TaqMan array card to detect the majority of known antimalarial resistance-associated mutations. The malaria TAC yielded excellent specificity, with no cross-reactivity to other pathogens or human genes and no unexplained cross-reactivity between alleles (Fig. S1 in the supplemental material; also data not shown). This tool displayed excellent sensitivity and accuracy when clonal laboratory parasites were the source of DNA and results were compared with results of Sanger sequencing (Tables 1 and 2). Additionally, the malaria TAC performed well when clinical samples were used (Tables 1 and 2), despite the presence of human DNA (~80 copies/reaction volume or 4.0 × 10^4 copies/100 μL of DNA). Based on the detection of heteroresistance and mixed genotypes, this tool has the ability to detect mixed infections (Table 3 and Fig. 2, S3, and S4), which are common, but underappreciated, in clinical samples.

Despite many benefits, there are also some obvious limitations of the malaria TAC. Although future malaria TAC designs are customizable, we are limited to the detection of known loci. For the most part, mutations that contribute to clinical resistance are well known, but new mutations whose significance remains unknown are still being discovered (37). Design flexibility is particularly relevant for emerging artemisinin resistance where novel Kelch13 resistance mutations are being detected (38). As is, our initial design of the malaria TAC appears to be a good approximation of important mutations. For example, we detected only two alleles in parasite DNA that were not included in the initial design (pfCRT 97L and pfDHPS 436F) (Table 1). These mutations were not originally represented because their global minor allele frequencies (MAF) were well below those for other alleles of these loci (39). Second, we detected ample diversity across the malaria TAC assays: (i) SNP barcodes were covered by 75% of
laboratory parasite lines and 61% of clinical samples, and (ii) resistance markers were covered by 81% of laboratory parasite lines and 38% of clinical samples.

From a genotyping standpoint, we were limited in the number of barcodes that could be interrogated. From just five loci, we detected distinct barcodes in 61% (41/67) of clinical samples. Since 19% of the barcodes were shared and since the majority of these were detected in at least two different countries, it is likely that an expansion of the number of barcodes tested would have revealed that many of these were indeed unique. For example, sample DB148 from Thailand had a genotyping barcode that was shared with two samples from China, three samples from Malawi, and one sample from Uganda. The malaria TAC also detected heteroresistance in this sample at two loci, indicating that it was likely a mixed infection. We propose that expansion of the number of SNP barcodes would have revealed a unique/mixed barcode in this sample that originated from Thailand. Future malaria TAC designs can augment this aspect of assay design.

A challenge to clinical infections is the detection of minor loci from mixed infections. Using the malaria TAC, we detected mixed infections in samples from all four countries. If we take into account both genotyping SNPs and observations of heteroresistance, Thailand exhibited 31%, China exhibited 10%, Malawi exhibited 33%, and Uganda exhibited 40% mixed infections. Of course, these were convenience samples that were not systematically collected for purposes of examining heteroresistance, so the absolute numbers are likely not representative of those regions. Four of seven cases of heteroresistance were also detected as mixed infections by genotyping SNPs, indicating that it is possible to accurately detect multiple alleles using the malaria TAC (down to 10% [data not shown]). Future assessment of the malaria TAC will work to accurately determine the limit of detection for rare alleles.

Clinical samples also exhibit various levels of parasite densities and thus parasite DNA. Measuring the limit of detection, or LOD, allows the estimation and comparison of sensitivity for various methods. Most of the malaria TAC assays displayed an LOD of

| Sample | SNP1 | SNP2 | SNP3 | SNP4 | SNP5 | SNP6 | SNP7 | SNP8 | Sample | SNP1 | SNP2 | SNP3 | SNP4 | SNP5 | SNP6 | SNP7 | SNP8 |
|--------|------|------|------|------|------|------|------|------|--------|------|------|------|------|------|------|------|------|
| DB123  | T    | A    | C    | C    | G    | G    | F11TC45 | T    | A    | C    | C    | G    | A    | IL5Z  | T    | A    | C    | C    | G    |
| DB201  | T    | A    | C    | C    | G    | G    | F11TC51 | T    | A    | C    | C    | A    | G    | 1LE0  | T    | A    | C    | C    | G    |
| DB139  | T    | A    | C    | C    | G    | A    | F11TC55 | T    | A    | C    | C    | A    | G    | 1CJ1  | T    | A    | C    | C    | A    |
| DB207  | T    | A    | C    | C    | G    | A    | F11TC10 | T    | A    | C    | G    | A    | G    | 0FSK  | T    | A    | C    | G    | G    |
| DB091  | T    | A    | C    | C    | A    | A    | F11TC14 | T    | A    | T    | C    | A    | A    | 090N  | T    | A    | C    | G    | G    |
| DB004  | T    | A    | C    | G    | G    | G    | F11TC31 | T    | A    | T    | C    | A    | A    | 0Z5G  | T    | A    | T    | C    | A    |
| DB076  | T    | A    | T    | G    | G    | A    | F11TC11 | T    | A    | T    | C    | G    | G    | 077R  | C    | A    | T    | C    | A    |
| DB002  | T    | G    | C    | G    | A    | A    | F11TC13 | T    | A    | T    | G    | G    | G    | 08K1  | C    | T    | C    | A    | A    |
| DB133  | C    | G    | C    | G    | G    | G    | F11TC05 | T    | G    | C    | G    | G    | A    | 0Z4T  | C    | A    | T    | C    | A    |
| DB085  | C    | G    | C    | G    | A    | A    | F11TC16 | T    | G    | C    | G    | A    | G    | 27LG  | C    | A    | C    | G    | A    |
| DB023  | C    | G    | T    | C    | A    | G    | F11TC07 | C    | A    | C    | C    | G    | G    | 01Y6  | C    | A    | C    | G    | A    |
| DB031  | C    | G    | T    | C    | A    | G    | F11TC17 | C    | A    | C    | C    | G    | G    | 1C9A  | T    | A    | T    | C    | G    | A    |
| DB055  | C    | G    | T    | C    | G    | G    | F11TC02 | C    | A    | C    | C    | G    | G    | 0ZDY  | C    | T    | A    | C    | G    | A    |
| DB043  | C    | A    | C    | C    | A    | A    | F11TC30 | C    | A    | C    | C    | G    | G    | 06KU  | C    | T    | A    | C    | G    | A    |
| DB107  | C    | A    | C    | C    | A    | A    | F11TC08 | C    | A    | C    | C    | A    | A    | 0HE3  | C    | A    | C    | G    | A    |
| DB111  | C    | A    | C    | C    | A    | A    | F11TC47 | C    | A    | C    | G    | G    | G    | C    | A    | C    | G    | A    |
| DB109  | C    | A    | C    | G    | A    | A    | F11TC09 | C    | A    | T    | C    | A    | A    | C    | A    | T    | C    | A    |
| DB206  | C    | A    | C    | G    | A    | A    | F11TC28 | C    | A    | T    | C    | A    | A    | C    | A    | T    | C    | A    |
| DB092  | C    | A    | T    | C    | A    | A    | F11TC01 | C    | A    | C    | C    | G    | A    | PI    | T    | A    | C    | C    | A    |
| DB148  | C    | A    | T    | C    | A    | G    | F11TC44 | C    | A    | C    | C    | G    | A/G  | U116  | T    | A    | T    | C    | A    |
| DB204RI| C    | A    | T    | C    | A    | G    | F11TC163| C    | A/G  | T/C  | C    | G    | G    | U113  | C    | A    | T    | C    | A    |
| DB203  | C    | A    | C    | C    | A    | A/G  | C    | A    | T    | C    | A    | A/G  | PI    | T    | A    | T    | C    | A    |
| DB206RI| C/T | A/T  | C    | G    | A/G  | C    | A/G  | T    | A/T  | C/G  | A    | A/G  | 165   | C    | A    | T/C  | C/G  | A/G  |
| DB104  | C/T | A/T  | C    | A    | A/G  | G    | C    | A    | T    | C    | A    | A/G  | 165   | C    | A    | T/C  | C/G  | A/G  |
| DB009  | T    | G    | T/C  | C    | G    | G    | C    | A    | T    | C    | A    | A/G  | 165   | C    | A    | T/C  | C/G  | A/G  |
| DB122  | T    | A    | T/C  | G    | A    | A    | C    | A    | T    | C    | A    | A/G  | 165   | C    | A    | T/C  | C/G  | A/G  |

Nonunique haplotypes are highlighted in the same color. Slashes indicate mixed strains (e.g., A/G). SNP1, Pf_01_000130573; SNP2, Pf_01_000539044; SNP3, Pf_02_000842803; SNP6, Pf_06_000145472; SNP7, Pf_06_000937750; SNP8, Pf_07_000277104.
40.3 copies/reaction volume (~8 to 40 parasites/μl, if single-copy genes were assessed on either the plate or TAC format). The malaria TAC LOD is similar to LODs of other TACs that we have developed (35). Additionally, the LOD is within range of other sensitive genotyping methods used in the malaria field (10 parasites/μl) (24, 40). Future work will explore the malaria TAC limit of detection using clinical samples.

One related concern that was revealed by these studies was the decrease in malaria TAC sensitivity when DNA derived from clinical parasite lines was used. This result is likely a limitation of the quality and quantity of the clinical samples that were used for this analysis. First, the amount of parasite-derived genomic DNA in these samples is unknown because of the presence of an abundance of human blood cell DNA. In order to correct for this, the maximum allowable sample volume was used when the clinical samples were used (20 to 50 μl of a 100-μl reaction mixture). Second, the format and age of the clinical samples varied, which could directly contribute to assay performance (see Materials and Methods for details). For example, the dried blood spots, remarkably, were collected 11 years ago in Thailand. Room temperature storage over long periods of time likely leads to some level of DNA degradation. Evidence for this was observed by the failure of amplification of long PCR product sizes suitable for Sanger sequencing (data not shown). Additionally, most of the undetectable samples were from Malawi; these purified DNA samples had the longest period (3 to 4 years) with an unknown number of freeze-thaw cycles. The cycle thresholds (Cₜ) of all malaria TAC assays provided evidence for a small amount/low quality of DNA in the Thailand and Malawi samples; for the hGAPDH assay, Uganda and China samples displayed average Cₜs of ~25 and 28, respectively (collected in 2008 to 2009 and 2016), while the Thailand (2005) and Malawi (2012 to 2013) samples had average Cₜs of ~32 and 34, respectively. These data indicate that freshly collected and prepared DNA would improve sensitivity of the malaria TAC.

While most assays on the malaria TAC performed very well, a few of the assays were problematic. pfCRT 72–76, pfCRT 97, pfDHFR 164, and pfDHPS 540 assays appeared suboptimal based on a sensitivity of <80% when they were tested with DNA from clinical parasites (Table 1). Three of these assays were the most challenging to design and exhibited the highest limits of detection. Conversely, pfCRT 97 performed well using laboratory parasite-derived DNA and exhibited an LOD similar to that of most of the other assays. It is possible that the presence of human DNA in clinical samples affects the performance of this assay. Although we did not specifically test the pfCRT 97 assay, we investigated this possibility by running five randomly chosen duplex assays in the presence and absence of human DNA at multiple ratios. Overall, we did not detect statistically significant differences during this analysis (data not shown). These results indicate that the presence of human DNA in clinical samples likely does not have an impact on these assays, but there is room for future work to investigate this further.

Although not the main purpose of this study, the malaria TAC detected a number of different resistance markers from several clinical isolates. For the most part, our observations are consistent with what has been observed previously. First, we did not detect pfCRT mutations from Malawi samples, consistent with the return of chloroquine-susceptible malaria in Malawi after chloroquine use was abandoned (41). The pfDHFR-pfDHPS mutant patterns IRNL-SGEGA and IRNL-AGEAA (mutations are underlined), which are linked to high-level sulfadoxine-pyrimethamine (SP) resistance (42), were observed as predominant alleles in Thailand and China samples, respectively. The pfMDR1 Y184F was the most common mutation observed in this study and has been reported as a common mutation found in Asia and Africa (43–45). Last, we found that 83% of pfKelch13 alleles from China harbored the F446I mutation; this is the predominant Kelch13 mutation associated with delayed clearance of parasites in patients close to the China-Myanmar border (9, 10).

Overall, our numbers of clinical samples from each country were small, and thus the sensitivity and specificity estimates of the malaria TAC are an approximation. It will be important to further evaluate this tool with additional clinical samples to fully explore its use for surveillance. Based on this pilot study of the malaria TAC, future design
alterations could include more genotyping loci (perhaps a new format), removal of cytochrome \( b \) mutations, and alteration of Kelch13 loci to better detect newly emerging resistance. Additionally, we can replace/redesign suboptimal assays (pfCRT 72–76, pfCRT 97, pfDHFR 164, and pfDHPS 540). Deployment of this molecular diagnostic technology requires an expensive real-time PCR platform (Viia7; Applied Biosystems); however, these instruments are already present in a number of countries of malaria endemicity, and this enables testing in country. We are hopeful that the malaria TAC will accelerate the ability to track antimalarial-resistant populations and impact local and national treatment recommendations.

**MATERIALS AND METHODS**

**Parasite and human DNA.** DNA from 16 *P. falciparum* lines with various mutations was used as controls in this study. Genomic DNA was directly obtained from BEI Resources (Manassas, VA, USA): *P. falciparum* 3D7 (MRA-102G), GB4 (MRA-925G), HB3 (MRA-155G), Dd2 (MRA-150G), 7G8 (MRA-152G), V1/S (MRA-176G), K1 (MRA-159G), TM90C6B (MRA-205G), W2 (MRA-157G), IpC 3445 (MRA-1236G), IPc 4884 (MRA-1238G), and 7C4/6 (MRA-172G). Cryopreserved parasites originally isolated from Battambang (MRA-1240 [BAT]), Mondolkriri (MRA-1241 [MON]), Pailin (MRA-1236 [PAIL]), and Pursat (MRA-1238 [PUR]), Cambodia, were also obtained from this laboratory, grown in our laboratory, and extracted for DNA as performed previously (46). Two non-*P. falciparum* species were also obtained for use as controls: genomic DNA of *P. knowlesi* (MRA-456G; BEI Resources) and cryopreserved *P. vivax* parasites (MRA-383; BEI Resources) from which DNA was directly extracted. Human control DNA was extracted from whole blood of healthy volunteers in our laboratory. All work was reviewed and approved by Institutional Biosafety and Human Investigation Committees at the University of Virginia.

**Plasmid controls.** Synthetic positive-control plasmids were constructed to cover resistance markers that were not represented in the above listed parasite lines (Genewiz, Inc., South Plainfield, NJ, USA). These loci included pfCRT25cVET, pfCRT25cVNT, pfCRT73–76VNT, pfCRT73–76VMT, pfCRT73–76VDT, pfCRT7Q, pfCYPt2858m, pfCYPt2868c, pfCYP2726n, pfKelch13–446l, pfKelch13–458y, pfKelch13–561h, pfKelch13–574l, pfKelch13–578s, and pfPlasmepsin4 of *P. malariae* and *P. ovale*. A synthetic positive-control plasmid was also constructed to contain the primer and probe regions of all 88 targets and used as a positive control for each malaria TAC run (Genewiz, Inc., South Plainfield, NJ, USA).

**Blood pathogens.** We tested the specificity of our malaria-specific assays against 35 other blood pathogens. Genomic DNAs from 11 pathogens were obtained from BEI Resources (Manassas, VA, USA), as follows: Acinetobacter baumannii (NR-10146), Brucella abortus (NR-2530), Haemophilus influenzae (DD-204), Klebsiella pneumoniae (NR-15465), Listeria monocytogenes (NR-13342), Pseudomonas aeruginosa (DD-723), Rickettsia sibirica (NR-10486), Staphylococcus aureus (HM-145D), Streptococcus pneumoniae (HM-145D), Toxoplasma gondii (NR-33509), and Yersinia pestis (NR-2720). Genomic DNAs for 15 pathogens were obtained from ATCC (Manassas, VA, USA), as follows: Bartonella bacilliformis (ATCC 35685D-S), Candida albicans (ATCC 10231D-S), Cryptococcus neoformans (ATCC 66031D-S), Leptospira interrogans (ATCC BAA-1198D-S), Leishmania infantum (ATCC 50134D), Neisseria meningitidis (ATCC 53415D-S), Streptococcus agalactiae (ATCC BAA-611D-S), Streptococcus pyogenes (ATCC 700294D-S), Enterococcus faecium (ATCC 51595D-S), Salmonella enterica serovar Typhi (ATCC 700931D-S), Staphylococcus epidermidis (ATCC 12228D-S), cytomegalovirus (ATCC VR-538D), Epstein-Barr virus (ATCC VR-3247D), herpes simplex virus 1 (ATCC VR-539DQ), and herpes simplex virus 2 (ATCC VR-540D). Genomic DNAs from nine Mycobacterium species were obtained from our collaborator (Department of Medicine Siriraj Hospital, Mahidol University, Bangkok, Thailand): *M. tuberculosis* ATCC 27294, *M. avium* ATCC 700898, *M. intracelle- lare* ATCC 13950, *M. kansasi* ATCC 12478, *M. simiae* ATCC 25275, *M. sherrisi* 553, *M. abscessus*, *M. fortuitum* ATCC 6841, and *M. peregrinum* ATCC 700868.

**Clinical samples and extractions.** Eighty-seven deidentified clinical samples were used to test the specificity of the malaria TAC. These included samples from Thailand (material, dried blood spots; collected 2005, \( n = 32 \)), China (material, purified DNA; collected 2016, \( n = 21 \) [47]), Malawi (material, purified DNA; collected 2012 to 2013, \( n = 28 \)), and Uganda (material, blood pellet or purified DNA; collected 2016, \( n = 1 \) [University of Virginia], and 2008 to 2009, \( n = 5 \) [48]). All clinical samples were collected as a part of previous studies that were reviewed and approved by Institutional Biosafety and Human Investigation Committees. When required, genomic DNA was extracted using QiAamp DNA minikit (Qiagen, Valencia, CA, USA) according to the manufacturer’s instructions and eluted in 100 μl of elution buffer.

**Confirmation by Sanger sequencing.** Mutations present in the control *P. falciparum* lines and a subset of clinical samples (42/87) were confirmed by Sanger sequencing. First, the resistance-associated genes and genotyping SNPs were PCR amplified using primers described in Table S5 in the supplemental material (pfCRT [chloroquine], pfCYP7 [atovaquone], pfDHFR [pyrimethamine], pfDHPS [sulfadoxine], pfMDR1 [multidrug resistant], pfKelch13 [artemisinin], Pf_01_000130573, Pf_01_000539044, Pf_02_000842803, Pf_06_000145472, Pf_06_000937750, and Pf_07_000277104). Each 25-μl PCR mixture contained 12.5 μl of HotStarTaq master mix (Qiagen, Valencia, CA, USA), 0.45 μl of the forward and reverse 50 μM primers (final concentration of 0.9 μM), 6.6 μl of nuclease-free water, and 5 μl of genomic DNA (500 pg total for DNA derived from laboratory parasites). PCR was performed on a CFX96 instrument (Bio-Rad, Hercules, CA, USA) and included an initial denaturation step at 95°C for 15 min, followed by 40 cycles of denaturation at 95°C for 30 s, annealing at 59°C for 30 s, and extension at 72°C for 30 s, with a final extension step at 72°C for 10 min. Next, PCR products were analyzed on 2% agarose gels, and
verified PCR products were purified using a MinElute UF PCR purification kit (Qiagen, Valencia, CA, USA) according to the manufacturer’s protocol. Finally, purified PCR products were measured spectrophotometrically, diluted with nuclease-free water, mixed with primers, and then submitted to GeneWiz (GeneWiz, Inc., South Plainfield, NJ, USA) for DNA sequencing. Sensitivity (the ability to correctly classify a mutant allele) and specificity (the ability to correctly classify a wild-type allele) were determined using a two-by-two table where the gold standard is Sanger sequencing. Accuracy (the ability to correctly classify both alleles) is an average of sensitivity and specificity values.

**Drug resistance loci and mutant allele selection.** We selected the targets for the initial malaria TAC design by (i) identifying previously reported associations with antimalarial resistance and (ii) prioritizing candidates based on global minor allele frequencies (MAF) reported in the MalariaGEN database (39). For *pfCRT*, *pfDHFR*, *pfDHPS*, and *pfMDR1*, most of the selected targets displayed a global MAF of >0.1 (5 of 26 total alleles) (Table S5). Since *pfKelch13* mutations have only recently arisen predominantly in Southeast Asia, global frequencies of all these are low (<0.1). We therefore included *pfKelch13* alleles that have been confirmed to confer artemisinin resistance (*Y493H, R539T, I543T, and CS80Y*) (14, 49) as well as one allele that is associated with clinically delayed parasite clearance (*F446I*) (10). Additionally, in order to provide a complete picture of variation across this gene, we included mutations that have been observed in Asia and Africa but at the time of selection had an unknown association with resistance (*N458Y, R561H, P574L, and A578S*) (50–52). Since data were not available for *pfCYTB* mutations in the MalariaGen database, we directly selected atovaquone resistance-associated alleles from the literature (18–20).

**Assay development in 384-well plate format.** Primers and TaqMan probes for the malaria TAC assays were either designed using Primer Express3 (44 of 87 assays; Applied Biosystems, Life Technologies Corporation, Carlsbad, CA, USA) or adopted from published sources (43 of 87 assays [23, 36, 53–59]) (Table 3A). Wild-type probes were designed based on the *P. falciparum* 3D7 sequences of the *pfCRT* (GenBank accession number NC_004328.2:458600–461695, gene ID 2655199), *pfCYTB* (AF069051.5), *pfDHFR* (NC_004318.1:755069–756895, gene ID 9221804), *pfDHPS* (NC_004329.2:549321–551737, gene ID 2655294), *pfMDR1* (NC_004326.1:1957885–962144, gene ID 813045), and *pfKelch13* (NC_004331.2:1724848–1727028, gene ID 814205) genes. Six primer sets/12 probes for genotyping SNPs, 5 primer sets/probes for species identification, and the human GAPDH internal control (hGAPDH) were described in previous publications (36, 57–59).

Optimization of conditions and probe specificity testing were performed using the 384-well plate on the ViiA7 platform (Applied Biosystems, Life Technologies Corporation, Carlsbad, CA, USA). Each assay was amplified in duplex (Fig. 1 shows pairings), except for assays involved in species identification and the 0.09 μM of each primer and 0.25 μM, respectively) were assayed in a 5-μl PCR mixture containing 2.5 μl of 2× TaqMan universal master mix II (Applied Biosystems, Life Technologies Corporation, Carlsbad, CA, USA), 1.27 μl of nuclease-free water, and 1 μl (1 ng/μl) of genomic DNA. Cycling conditions included an initial denaturation at 95°C for 10 min, followed by 40 cycles of denaturation at 95°C for 15 s and annealing/extension at 59°C for 1 min. DNA sources included either *P. falciparum* 3D7 DNA as a wild-type control, 10 sequence-confirmed mutant *P. falciparum* lines (including Dd2, V1/S, 7G8, TMC90C6B, K1, HB3, PAIL, PUR, MON, and BAT), *P. vivax*, *P. knowlesi*, synthetic mutant control plasmids and species identification plasmids (*P. malariae* and *P. ovale*), a mixture of parasite-human DNA to test the performance of assays in the presence of human DNA (see below for ratios), or nuclease-free water as a nontemplate control.

Assay amplification efficiency and limits of detection were first performed on the 384-well plate format and subsequently on the array card format. To do so, DNA from individual parasite lines or plasmid controls was 10-fold serially diluted (plasmid control range, 105 to 1 copy/nl; parasite DNA range, 1 ng to 10 fg or the equivalent of 4.03 x 104 to 0.403 copies/μl). For 384-well plate assays, 1 μl of diluted samples was tested in each 5-μl reaction mixture in triplicate. Since the volume of DNA used in the array card is 5-fold lower (0.2 μl reaction mixture), dilutions for malaria TAC testing were prepared as 5-fold more concentrated to ensure equivalence on both formats. The copy number of plasmid controls and copy number equivalents for parasite DNA were calculated using an available online tool (University of Rhode Island Genomics and Sequencing Center calculator [http://cels.uri.edu/gsc/cndna.html]) by inputting the amount of DNA in nanograms and the length in base pairs (23 Kb was used for parasite DNA). The formula is as follows: number of copies = (amount of DNA x 6.022 x 1024)/(length of template x 1 x 104 x 650). To determine whether human DNA in the clinical samples impacted the performance of our assays, five randomly chosen assays were tested in duplicate in the presence and absence of human DNA (*pfCRT* 72C/725tc, *pfDHFR* 164I/164L, *pfDHPS* 540K/540E, *pfMDR1* 86N/86Y, and *P. falciparum*). This was performed at multiple ratios of parasite/human DNA; human DNA was fixed at 10 ng (2,860 genome copies), and parasite DNA varied from 1 ng to 1 pg (40,300 to 40.3 genome copies), yielding the following ratios: 14:1, 1:4:1, 0.14:1, and 0.014:1.

**Evaluation of the malaria TAC.** Primer and TaqMan probe oligonucleotides were custom ordered, synthesized, and spotted into the microfluidic card by Applied Biosystems (Life Technologies Corporation, Carlsbad, CA, USA) as laid out in Fig. 1. Twenty to 50 μl of input DNA (at 1 ng/μl for culture parasite-derived DNA) was mixed with 50 μl of 2× TaqMan universal master mix II (Applied Biosystems, Life Technologies Corporation, Carlsbad, CA, USA) and 0 to 30 μl of nuclease-free water to yield a 100-μl final volume. This mixture was loaded into each port of the card; each card included a port for seven clinical samples and one synthetic positive-control plasmid (8 ports total). The loaded card was centrifuged twice at 1,200 rpm for 1 min and then sealed. The loading ports were excised, and the full card was inserted into a ViiA7 instrument (Life Technologies Corporation, Carlsbad, CA, USA) and run under
the same cycling conditions as described above for 45 cycles. The results were automatically analyzed by Viia 7 software; the baseline and threshold were adjusted in cases where minor cross-reactivity occurred.

Statistical analysis. Means or medians were compared using Student’s t test or a Mann-Whitney test. Data are shown as means ± standard deviations unless otherwise stated. A standard curve of hGAPDH was generated with known DNA concentrations and plotted against the Cₜ value to yield the following equation: copy number per reaction product = 10^{-H11002/37.65}/H11002/37.65 + 1.544.

SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at https://doi.org/10.1128/AAC.00110-17.

SUPPLEMENTAL FILE 1, PDF file, 2.0 MB.

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