Complex DNA structures trigger copy number variation across the Plasmodium falciparum genome

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ABSTRACT

Antimalarial resistance is a major obstacle in the eradication of the human malaria parasite, Plasmodium falciparum. Genome amplifications, a type of DNA copy number variation (CNV), facilitate over-expression of drug targets and contribute to parasite survival. Long monomeric A/T tracks are found at the breakpoints of many Plasmodium resistance-conferring CNVs. We hypothesize that other proximal sequence features, such as DNA hairpins, act with A/T tracks to trigger CNV formation. By adapting a sequence analysis pipeline to investigate previously reported CNVs, we identified breakpoints in 35 parasite clones with near single base-pair resolution. Using parental genome sequence, we predicted the formation of stable hairpins within close proximity to all future breakpoint locations. Especially stable hairpins were predicted to form near five shared breakpoints, establishing that the initiating event could have occurred at these sites. Further in-depth analyses defined characteristics of these ‘trigger sites’ across the genome and detected signatures of error-prone repair pathways at the breakpoints. We propose that these two genomic signals form the initial lesion (hairpins) and facilitate microhomology-mediated repair (A/T tracks) that lead to CNV formation across this highly repetitive genome. Targeting these repair pathways in P. falciparum may be used to block adaptation to antimalarial drugs.

INTRODUCTION

Major efforts have succeeded in eradicating malaria in North America and Europe, but have largely failed in Southeast Asia and Africa (1). Some of the remaining challenges include a lack of accessible treatments and the widespread development of drug resistance. Plasmodium falciparum, the protozoan parasite that causes the most severe form of malaria and the majority of malaria deaths, has developed resistance to all drug interventions thus far (2). Single nucleotide polymorphisms (SNPs) are the most commonly studied genetic contribution to antimalarial drug resistance. However, chromosomal size polymorphisms, including copy number variations (CNVs) that encompass the genes of antimalarial targets or drug transporters, also play a key role in parasite survival (3).

CNVs often carry strong fitness costs due to increased cellular burden for DNA replication and alterations of metabolic flux due to differing levels of enzyme expression (4). However, it has been proposed that in many organisms, including P. falciparum, the creation of redundant gene copies facilitates the accumulation of SNPs (5–8). Studies observing both types of mutations in Plasmodium provide evidence that CNVs appear to eventually be lost in favor of SNPs (9–11).

Two CNVs associated with clinical antimalarial resistance encompass the genes encoding the multiple drug resistance protein 1 (pfmdr1) and GTP-cyclohydrolase 1 (gch1) (12–17). Additionally, a number of resistance-associated CNVs across many chromosomes were detected in the P. falciparum genome following laboratory selections with novel antimalarials (8,9,13,16,18–25). CNVs have also been detected in clinical Plasmodium vivax isolates (18,26–29), providing evidence that this form of adaptation is not confined to P. falciparum.

Mechanisms leading to CNVs in Plasmodium are currently unknown. Due to a lack of significant sequence homology surrounding the CNV breakpoints, homologous recombination is not likely to be involved in the process. The most compelling evidence of a shared mechanism is the presence of long monomeric A/T tracks at CNV bound-
aries (8,17,27,30,31). In other organisms, there is precedence for polymerase pausing and DNA double-stranded breaks (DSBs) at long mononucleotide repeats or AT/TA dinucleotide repeats (32–35). However, in-depth characterization of multiple independently generated CNVs on chromosome 6 indicates an additional signal present that triggers amplification (8). Specifically, two distinct CNVs were found to share a common boundary on one end (C and F clones, Figure 1A), an event that is highly unlikely to occur by chance. The A/T track at this shared breakpoint is not significantly longer (37 bp long compared to a mean of 33 bp for all CNVs included in our analysis) and thus other factors must be driving this repeat occurrence. Abnormal DNA structures, including hairpins and stem-loops, have also been implicated in replication fork stalling and DSBs in yeast and humans (36–41). Therefore, we investigated whether sequences proximal to CNV breakpoints across the highly A/T-rich P. falciparum genome are enriched in these DNA structures.

Here, we present evidence that DNA hairpin formation is likely an initiating event in the generation of CNVs in P. falciparum. First, we adapted a CNV-calling pipeline to achieve near single base pair resolution to study laboratory acquired CNVs in 35 total resistant parasite clones selected with eight different antimalarials (19 parasite clones described (50,51). In brief, 50 bp windows were selected by 3′ and downstream (3′) from the ψ end) and upstream (5′ end) from the pre-CNV, see below). For clones in which whole genome sequencing was not available (DSM1E and F), previously published sequence from PCR amplification across the breakpoint was used to pinpoint precise breakpoint locations (8).

MATERIALS AND METHODS
Collecting genome and breakpoint sequences
We analyzed whole genome sequencing data to identify CNVs from in vitro haploid erythrocytic P. falciparum parasites that were selected with a number of different antimalarial clones (Table 1, see details on parent and resistant clones, antimalarial target, chromosome, CNV sizes and accession numbers in Supplementary Table S1 (8,16,42,43)). For clarity of procedures, we present a flow chart of our overall analysis methods (Supplementary Figure S1). Briefly, low quality bases and adapter sequences from Illumina-based whole genome sequencing of both the parasite during adaptation to countless antimalarials. These ‘trigger sites’ are found broadly throughout the parasite genome and would facilitate adaptation to most selective forces. In-depth analysis of breakpoints in resistant clones (termed post-CNВ) suggests the action of two repair pathways that utilize the A/T tracks as short stretches of homology. These findings contribute to a growing model of the mechanisms that lead to enhanced generation of CNVs across highly repetitive genomes.

Calculating the likelihood of DNA hairpin formation
The probability of hairpin structure formation across the desired regions was predicted essentially as previously described (50,51). In brief, 50 bp windows were selected by shifting by 1 bp across a 2 kb stretch of sequence surrounding the pre-CNВ breakpoint position in the parent genome. The 50 bp windows were chosen to ensure hairpin formation was possible within the Okazaki initiation zone during
Figure 1. Highly stable DNA hairpins are found near pre-CNV boundaries. Resistant clones from various selections exhibit a range of CNV sizes but all have long A/T track breakpoints on their upstream and downstream ends (see Supplementary Table S3). Shared breakpoints are indicated with arrows and depicted in red (boxes and plots); unique breakpoints are shown for comparison and depicted in gray (boxes and plots). Although 2kb was analyzed, for simplicity, the insets show the ΔG of folding for each 50 bp window across 1 kb of sequence surrounding the A/T track breakpoint (vertical red/gray bar at 500 bp). The dotted line demarks the threshold for stable hairpin formation (ΔG of −5.8 kCal/mol, see ‘Materials and Methods’ section for how this was defined). (A) Each CNV from DSM1 resistant parasites (C, D, E and F) encompasses the gene for the target dihydroorotate dehydrogenase (DHODH, gray bar with star). The shared 3′ breakpoint from clones C and F is indicated (arrows). (B) Each CNV from Cladosporin resistant parasites (A, B and C) encompasses the gene for the target lysyl-tRNA synthetase (KRS1, gray bar with star). The shared 3′ breakpoint from CladoA and CladoC is indicated (arrows). (C) Each CNV from the MMV019662 and MMV028038 resistant parasites (1F4, 2G6, 3B6 and 2B6) encompasses the gene target Pf3D7_0107500, a member of the resistance-nodulation-division transporter family (gray bar with star). The shared breakpoints are indicated (arrows).
Replication. The size of the Okazaki initiation zone is not known in *Plasmodium* but it is expected to be in the same range as other eukaryotes (300–1000 bp (52)). Next, the Gibbs free energy (ΔG), which predicts the stability of the sequence folding on itself, was determined for each window using Vienna 2.1.9 folding prediction software with Mathews 2004 DNA folding parameters and G-quadruplexes, GU pairing, and lonely base pairs were disallowed (53). Lonely base pairs are helices in a hairpin or stem-loop that are composed of only 1 bp and do not stack on other base pairs. These structures are not energetically favorable and cannot form and are therefore excluded from analyses. During this analysis, each 50 bp window was counted as a separate possible hairpin. Initially this analysis was confined to sequences from the parent genome prior to CNV generation (the pre-CNVT breakpoint position). Predictions were subsequently performed on sequences from post-CNVT breakpoint localizations from resistant clones.

### Defining stable hairpins

Due to a non-normal distribution of predicted hairpin ΔG-values, the ΔG cut-off of stable hairpins was determined using a randomization method: sequence from each chromosome was randomly shuffled using the EMBOSS shufflesseq function to maintain overall A/T composition and hairpins were again predicted (54). In this analysis, 50 kb of sequence on either chromosome end was trimmed to avoid highly repetitive telomeric sequences. The value of the resulting top 3% of shuffled hairpins was used as the stability cut-off for all analyses (−5.8 kCal/mol); sequences with values below this cut-off indicated a high probability of a ‘stable’ structure forming. This value is consistent with that utilized in previous *P. falciparum* investigations (51). Furthermore, this value is similar to the top 5% of non-shuffled hairpins (ΔG of −5.5 kcal/mol in our analysis), a threshold utilized in secondary structure studies of other organisms (55).

### Determining the mean ΔG profile

The mean ΔG of folding in close proximity to CNV breakpoints (shared or all) was determined by setting the end of the A/T track breakpoint to distance zero and calculating the mean ΔG for each 50 bp window as the sequence is shifted by 1 bp. The 95% confidence interval of each position was calculated and then plotted using Graphpad PRISM 7 (www.graphpad.com). For comparison with sequences not associated with CNVs, this process was repeated with 36 randomly chosen A/T tracks across the genome (Supplementary Table S4 and see ‘Evaluating A/T track lengths across the genome’ section). Each random A/T track position was chosen using a random number generator to pick a line number from the bed file of all A/T tracks of this size across the genome (excluding telomeres). Due to unequal sample sizes and a non-normal distribution, the level of significance in differences was calculated using the Wilcoxon–Mann–Whitney test.

### Evaluating A/T track lengths across the genome

A/T tracks were identified with the Phobos Repeat Finder (Version 3.3.11, http://www.rub.de/ecoevo/cm/cm.phobos.htm), which mapped the locations and lengths of long monomeric A/T tracks > 9 bp across the 3d7 genome (Supplementary Figure S1C). The level of significance in differences between the two datasets was again calculated using Wilcoxon–Mann–Whitney test. This length of track was chosen based on a previous study that showed that those

### Table 1. Characteristics of *Plasmodium falciparum* CNVs used in this study

| Clone       | Data source         | CNV Chr. | CNV start (bp w/ 95% confidence interval) | CNV end (bp w/ 95% confidence interval) | # of supporting genomes |
|-------------|---------------------|----------|------------------------------------------|----------------------------------------|-------------------------|
| DSM1C       | Guler et al., (8)   | 6        | 79 067 ± 0                               | 152 482 ± 0                            | 1                       |
| DSM1D       |                     | 6        | 64 578 ± 0                               | 158 152 ± 0                            | 1                       |
| DSM1E       |                     | 6        | 118 425 ± 0                              | 153 231 ± 0                            | 1                       |
| DSM1F       |                     | 6        | 113 523 ± 0                              | 152 482 ± 0                            | 1                       |
| HFGRII      | Herman et al., (42) | 12       | 587 623 ± 61                             | 612 922 ± 3                            | 1                       |
| HFGRIII     |                     | 12       | 589 189 ± 5                              | 621 909 ± 1                            | 1                       |
| CladoA      | Manary et al., (60) | 13       | 2 000 221 ± 11                           | 2 058 842 ± 1                          | 1                       |
| CladoB      |                     | 13       | 2 004 915 ± 2                            | 2 055 159 ± 1                          | 1                       |
| CladoC      |                     | 13       | 2 000 213 ± 4                            | 2 022 803 ± 0                          | 1                       |
| PQA11       | Cowell et al., (43) | 10       | 290 655 ± 0                              | 308 771 ± 2                            | 1                       |
| F7          |                     | 1        | 264 317 ± 0                              | 359 349 ± 0                            | 6                       |
| 3B6         |                     | 1        | 264 317 ± 1                              | 362 912 ± 0                            | 1                       |
| 1F4         |                     | 1        | 321 511 ± 5                              | 373 058 ± 9                            | 2                       |
| 2G9         |                     | 1        | 321 511 ± 2                              | 362 912 ± 10                           | 2                       |
| 1E3         |                     | 1        | 321 511 ± 2                              | 362 913 ± 9                            | 2                       |
| 3NXC3       |                     | 12       | 1 733 591 ± 3                            | 1 768 713 ± 3                          | 1                       |
| 3C3         |                     | 12       | 1 718 154 ± 3                            | 1 769 038 ± 3                          | 6                       |
| R2B2        |                     | 3        | 782 909 ± 45                             | 845 526 ± 0                            | 4                       |
| 1B2 ch10    |                     | 10       | 285 731 ± 27                             | 315 681 ± 3                            | 1                       |
| 1B2 ch12    |                     | 12       | 1 549 855 ± 5                            | 1 567 426 ± 1                          | 1                       |

*Whole genome sequencing is not available for these two clones. Analysis was performed using locations identified by PCR sequencing.*
above 9 bp were over-represented on *P. falciparum* chromosome 2 (56). To determine if A/T tracks were observed solely due to the high A/T content of *P. falciparum* (80.6%), we calculated the probability of observing different A/T tracks lengths based purely on nucleotide composition. Frequencies of monomeric A/T tracks of length \( N \) were calculated as follows:

The observed frequency of \( A \) and \( T \) tracks of length \( N \) were obtained using the following equation:

\[
f_N^{\text{obs}} = \frac{C_N^{\text{obs}}}{l_{\text{seq}}}
\]

Where \( C_N^{\text{obs}} \) is the observed number of monomeric tracks of length \( N \) and \( l_{\text{seq}} \) is the length of the chromosome sequence.

For each \( A \) or \( T \) track observed with length \( N \), the corresponding expected frequency of mononucleotide \( A \) and \( T \) tracks was obtained from the following equation:

\[
f_N^{\text{exp}} = \left( f_A^{\text{obs}} \right)^N \left( 1 - f_A^{\text{obs}} \right)^2 + \left( f_T^{\text{obs}} \right)^N \left( 1 - f_T^{\text{obs}} \right)^2
\]

where \( f_A^{\text{obs}} \) is the observed frequency of any base pair which corresponds to the overall percent base composition.

Maximum expected length for each chromosome was found using the following formula:

\[
N_{\text{exp}} = \frac{\log \left( \frac{1}{\log \left( f_A^{\text{obs}} \right)} \right)}{\log \left( f_A^{\text{obs}} \right)} + \frac{\log \left( \frac{1}{\log \left( f_T^{\text{obs}} \right)} \right)}{\log \left( f_T^{\text{obs}} \right)}
\]

**Investigating genome scale A/T track-hairpin relationships**

In order to assess the hairpin and A/T track relationship on a larger scale, hairpins across the entire genome were predicted as described above. Where indicated, analyses were confined to tracks >20 bp as this reflects the lengths of A/T tracks found at observed CNV breakpoints (Supplementary Table S4). The relationship between hairpins and long A/T tracks was then determined in genic and intergenic regions separately. This was accomplished by taking gene annotations from the 3d7 reference genome and extracting A/T tracks from regions within or outside of gene annotations utilizing the 'intersect' and 'subtract' bedtools functions, respectively (Supplementary Figure S1C). Distance between genic or intergenic A/T tracks to the nearest stable hairpin (either upstream or downstream) was then calculated using the 'closest' function in bedtools (57). For this analysis, the positions of the local minima of hairpins had to be identified. First, we extracted all hairpins below our significance threshold (−5.8 kcal/mol, see ‘Defining stable hairpins’ section). Then, for each set of windows with contiguous positions below this threshold, we identified the window with the most negative value and created a data subset with these minima. If there were multiple contiguous windows with the same value, all matching windows were extracted and used for analysis. The level of significance in differences was calculated using the Wilcoxon–Mann–Whitney test. Visualization of the frequency of lengths of the A/T tracks compared to the distance to stable hairpins was performed using ggplot2 in R version 3.2.4 (58,59). The Kolmogorov–Smirnov non-parametric test was used to compare the equality of intergenic and genic distributions to determine significant differences.

**RESULTS**

**CNV breakpoint features are conserved in *Plasmodium falciparum***

We obtained sequence from *P. falciparum* clones that had been selected for resistance to novel antimalarials in vitro (8,16,43,60) (Supplementary Table S1). After read alignment and CNV calling using an adapted Speedseq pipeline with stringent quality controls (see ‘Materials and Methods’ section), we selected sequence from 35 parasite clones that displayed high confidence CNV breakpoints for further analysis (Table 1 and Supplementary Table S2). Due to improved resolution, breakpoint locations were identified primarily through discordant- and split-read analysis extracted by LUMPY. This analysis identified 19 distinct CNVs for a total of 33 CNV breakpoints: 5 were conserved between different CNVs in multiple parasite clones (termed ‘shared’ breakpoints) and 28 were unique to their respective CNV (Table 1). In total, these breakpoints had a median of 27 supporting split and discordant reads (range of 3–1025 reads, Supplementary Table S3). Read depth changes detected by CNVnator further confirmed these general breakpoint locations and the orientation of reads confirmed the tandem duplications at these sites (Supplementary Figure S2).

Confidence in this analysis was bolstered by overall read depth and quality scores determined for each sequenced genome. Read depth across each chromosome, excluding telomeric regions, was >40-fold (median of 87-fold); coverage across CNV breakpoints was similar with a median of 107-fold for 2 kb surrounding breakpoints and a median of 57-fold for 100 bp surrounding breakpoints (Supplementary Table S2). The mean mapping quality scores across the genome was 57 out of a maximum score of 60 (44).

To determine whether DNA hairpins were associated with CNV breakpoints in *P. falciparum*, we went to the locations of the shared breakpoints in the pre-CNV parent genome. Two kilobases of proximal sequence were used to predict the probability of secondary structure formation nearby; a \( \Delta G \) of \(<−5.8\) kcal/mol indicated a high probability of a ‘stable’ structure forming from this sequence window (see ‘Materials and Methods’ section). From this focused analysis, we invariably detected extremely stable hairpins (the top 0.2% most stable structures across the entire genome, mean \( \Delta G \) of \(<−9.7\) kcal/mol) within a few hundred base pairs of the shared breakpoint A/T tracks (Figure 1A–C, mean distance of 165 ± 58 bp, Table 2). Stable hairpin structures were predicted to form by inverted repeats and AT dinucleotides present in the analyzed sequence (Table 2).

In all cases, multiple stable hairpins were detected in close proximity to the shared breakpoints (see Figure 1, where multiple peaks reach or fall below the dotted line); it was not clear which structure was contributing to CNV formation, the closest or the most stable hairpin. We therefore used this data to investigate whether there was a critical A/T track-hairpin distance; we determined the mean \( \Delta G \) at each
Table 2. Hairpin stability and distance relationships at CNV breakpoints

| Breakpoint       | ΔG of closest hairpin | Track-hairpin distancea | Hairpin forming sequence |
|------------------|-----------------------|--------------------------|--------------------------|
| DSM1F/C/3        | −10.9                 | 88                       | Inverted repeat          |
| CladoA/C/5       | −9.1                  | 222                      | Inverted repeat          |
| F7/3B6.5         | −8.1                  | 218                      | AT dinucleotide          |
| 1F4/1E3.5        | −10.2                 | 104                      | AT dinucleotide          |
| 3B6/1E3.3        | −10.2                 | 194                      | AT dinucleotide          |
| Mean of shared   | −9.7 ± 1.0            | 165 ± 58                 | NA                       |
| DSM1C/5          | −6.7                  | 216                      | Inverted repeat          |
| DSM1D.5          | −9.7                  | 49                       | Inverted repeat          |
| DSM1D.3          | −7.1                  | 2                        | Inverted repeat          |
| DSM1E.2b         | −6.3                  | 234                      | AT dinucleotide          |
| DSM1E.2b         | −5.8                  | 424                      | Inverted repeat          |
| DSM1F.5b         | −8.4                  | 172                      | AT dinucleotide          |
| HFGRII.5         | −6.1                  | 0                        | Inverted repeat          |
| HFGRII.3         | −7.3                  | 2                        | Inverted repeat          |
| HFGRIII.5        | −7.1                  | 21                       | Inverted repeat          |
| HFGRIII.3        | −6.7                  | 137                      | AT dinucleotide          |
| CladoA.3         | −7.9                  | 59                       | AT dinucleotide          |
| CladoB5          | −6.6                  | 105                      | Inverted repeat          |
| CladoB3          | −6.1                  | 14                       | AT dinucleotide          |
| CladoC3          | −8.3                  | 92                       | AT dinucleotide          |
| PQA11.5          | −13.2                 | 234                      | AT dinucleotide          |
| PQA11.3          | −8.7                  | 212                      | AT dinucleotide          |
| 1F4.3            | −8.9                  | 268                      | AT dinucleotide          |
| 2G9.5            | −13.1                 | 104                      | AT dinucleotide          |
| 2G9.3            | −10.2                 | 194                      | AT dinucleotide          |
| 33X3C.5c         | −13.1                 | 118                      | AT dinucleotide          |
| 33X3C.3c         | −9                    | 30                       | Inverted repeat          |
| 3C3              | −6.8                  | 342                      | Inverted repeat          |
| 3C3              | −9                    | 261                      | Inverted repeat          |
| R2B2.5           | −7.3                  | 2                        | Inverted repeat          |
| R2B2.3           | −10.7                 | 95                       | AT dinucleotide          |
| 1B2ch10.5        | −8.4                  | 0                        | AT dinucleotide          |
| 1B2ch10.3        | −8.3                  | 123                      | AT dinucleotide          |
| 1B2ch12.5c       | −8                    | 2                        | AT dinucleotide          |
| 1B2ch12.3c       | −7.9                  | 171                      | AT dinucleotide          |
| Mean of all      | −8.6 ± 2.0            | 132.6 ± 106.3            | NA                       |

NA, not applicable. _5_, upstream breakpoint. _3_, downstream breakpoint.

aTrack-hairpin distance was calculated to the nearest stably predicted hairpin. Distances of 0: the A/T track breakpoint is participating in hairpin formation.

bSequences derived from PCR across breakpoints.

cUtilize A/T dinucleotides as the breakpoint rather than A/T tracks.

base pair traveling away from the A/T tracks for the five shared breakpoints. When we compared this profile with that from random A/T tracks across the genome that do not participate in CNV formation (see ‘Materials and Methods’ section for details about these sequences were chosen), we detected a ΔG minima for the shared breakpoints at a distance of ~80 and ~360 bp (Figure 2A, P < 0.05 for both). This analysis provided evidence that stable hairpins within very close proximity, <400 bp, to the breakpoint A/T track likely contributed to CNV formation.

We extended our analysis to the remainder of the high-quality CNV breakpoints identified in the above analysis (Supplementary Table S1). Although less pronounced than with the shared breakpoints, the mean ΔG profile for all CNV breakpoints indicated that the most stable structure is within ~400 bp (Figure 2B). Minima were identified at similar distances from the breakpoints and were significantly stronger than random A/T tracks (P < 0.05). In line with this result, stably predicted hairpins were found in very close proximity to all CNV breakpoints (mean hairpin distance of 133 ± 106 bp, mean ΔG of −8.6 kCal/mol). Overall, 42% of

Figure 2. Mean-free energy profiles highlight a critical distance for stable hairpins. The mean ΔG of folding in close proximity to shared (A) and all (B) CNV breakpoints is plotted. This was done by setting the A/T track breakpoint at a distance of 0 bp and calculating the mean ΔG for each window of 50 bp as the sequence is shifted by 1 bp (A: purple line and B: blue line). As a comparator, the mean ΔG profile of 36 randomly chosen A/T tracks not associated with CNV formation (20–40 bp in length) was plotted (green line, see characteristics in ‘Materials and Methods’ section). Mean values with 95% confidence interval are shown. Shared breakpoints are DSM1C/F.3, CladoA/C.5, F7/3B6.5, 1F4/1E3.5 and 3B6/1E3.3 (see Table 2)
breakpoints had a highly stable structure within 100 bp of the A/T track breakpoint, 60% within 150 bp distance and all but one within 400 bp (Table 2). These proximal structures were frequently composed of inverted repeats or AT dinucleotide repeats (Table 2).

As has been noted before, the majority of CNV breakpoints occurred at very long A/T tracks (>20 bp, Supplementary Table S4). There were a few exceptions; AT dinucleotide repeats sat at both junctions for 33XC3 and 1B2 ch10 and an imperfect A/T track was found on the 3’ end of the 1F4 clone (88% pure T’s).

**CNV breakpoint features are enriched in intergenic regions**

We noted previously that CNV breakpoints are more often found in intergenic than genic regions (8). To explore this further, we expanded our analysis across these two regions of the *P. falciparum* genome. Specifically, we investigated (i) the quantity and length of A/T tracks, (ii) the propensity for DNA hairpin formation, as measured by ΔG of folding and (iii) the distance relationship between these two features. First, when compared to expected numbers, long A/T tracks >9 bp were highly enriched across the genome (Supplementary Figure S3, *P* < 0.01 for A/T tracks > 9 bp). When comparing genic to intergenic regions of the genome, we found about twice as many long A/T tracks in intergenic sequences than genic (42 026 in intergenic versus 19 408 in genic, Table 3, *P* < 0.001). A more striking difference was observed if the quantity of very long A/T tracks, >20 bp, were compared (~4-fold increase: 9509 in intergenic regions and 2410 in genic, *P* < 0.001). Second, we predicted a greater number of stable structures (ΔG < −5.8 kCal/mol) in intergenic compared to genic regions (37 439 intergenic and 23 442 genic, Table 3, *P* < 0.05) and an increase in the mean hairpin strength of these stable hairpins (~7.56 kCal/mol for intergenic compared to −7.23 kCal/mol for genic, *P* < 0.01). Finally, we found that the distance between A/T tracks and hairpins differed greatly between genic and intergenic regions. The mean A/T track-hairpin distance when considering long A/T tracks was 99 bp in intergenic regions and 277 bp in genic regions (Table 3, *P* < 10⁻¹³). This trend was conserved when considering very long A/T tracks (mean of 104 bp distance in intergenic and 163 bp in genic, *P* < 10⁻⁶).

By visualizing these distributions on a whole genome scale, the disparities between the two genomic regions and the close A/T track-hairpin association in intergenic regions are emphasized (Figure 3A and B, Kolmogorov–Smirnov test, *P* < 10⁻¹⁵). Due to the characteristics of these features that are associated with observed CNV breakpoints, we propose that there is an optimal range for A/T track lengths (~20–40 bp) and track-hairpin distances (<400 bp) (yellow highlight in Figure 3A and B). We defined genome positions with these characteristics as CNV ‘trigger sites’: those locations that are competent to generate CNVs. Using these parameters, there are 9130 intergenic and 2222 genic trigger sites across the *P. falciparum* genome which corresponds to 19.0% of intergenic and 9.6% of genic A/T tracks.

**Identifying DNA repair pathways utilized in CNV formation**

The above analysis was performed using parent sequence **prior to** CNV formation (pre-CNV, Figure 4A). In order to pinpoint which repair pathways may be acting in this process, we also studied the sequence from resistant clones **after** CNV formation (post-CNV, Figure 4B). This was accomplished by comparing pre- and post-CNV sequences from two sources, when available: PCR sequence of the A/T track breakpoint (for two DSM1 resistant clones) and split-reads from breakpoint alignment sequences (for another 14 clones). We found that the post-CNV A/T track lengths were 16.6 ± 19.0% shorter than the pre-CNV lengths (Supplementary Table S4, *P* < 0.01). Despite the almost ubiquitous shortening of the breakpoint A/T track, hairpin predictions using post-CNV sequence from DSM1 resistant clones yielded a pattern similar to that of pre-CNV sequence due to a general lack of mutations surrounding the A/T tracks (Figure 4C; Supplementary Figure S4B and D). In two exceptions (of seven post-CNV breakpoints analyzed), a novel stable hairpin was generated (Figure 4D; Supplementary Figure S4A and C), indicating sequence changes following CNV generation. Analysis of deep sequencing reads at these locations further confirmed these findings (unpublished data). These two different patterns suggest the action of multiple repair pathways in CNV generation (see ‘Discussion’ section).
shared CNV breakpoints reveal a model of CNV formation

High-quality deep sequencing of parasites from several controlled laboratory selections provided a unique opportunity to study CNV formation in the *P. falciparum* genome (Supplementary Table S1). Three characteristics facilitated these studies: (i) the availability of sequence from parent clones (prior to selection or pre-CNV) allowed for analysis of the native genome architecture at the position of the future CNV breakpoint, (ii) sequence from resistant clones (post-CNV) allowed for mechanistic studies on the pathways that enacted the change and (iii) breakpoints that occurred more than once in independent selections (or ‘shared’ breakpoints) allowed us to identify features that likely contribute to CNV formation.

Overall, five shared breakpoints were detected in our analysis; due to their occurrence, we speculated that there was an additional CNV signal beyond the almost ubiquitous A/T track present at these locations. Indeed, secondary structure predictions identified extremely stable hairpins in close proximity to these shared breakpoints (Figures 1, 2 and Table 2). The specific hairpins identified in this analysis were more stable than 99.8% of hairpins predicted across the genome (∼23.5 million structures overall) or the top 8% of stable hairpins (∼61 000 structures with ΔG of −5.8 kcal/mol in total). This finding increased our confidence that hairpins within close proximity to the breakpoint A/T track were of importance. Structure predictions on the remaining unique CNV breakpoints displayed a similar profile with a mean ΔG in the top 12% of stably predicted hairpins across the genome.

DNA hairpins and other secondary structures have been implicated in mechanisms of immune evasion by *P. falciparum* (41, 51, 73). Additionally, such structures are known to cause problems during DNA replication in other organisms: they result in higher levels of replication fork collapse and DNA breakage (40, 74) and hairpin-binding proteins can stimulate recombination at these sites (75–77). When repaired erroneously, these events can lead to the formation of CNVs (50, 51, 74, 78).

In light of these previous studies and our results, we propose a model of CNV generation (Figure 5): DNA hairpins in close proximity to long A/T tracks throughout the *P. falciparum* genome have the propensity to create DSBs by...
Resulting amplifications are initially rare throughout the genome and are repaired in a non-faithful manner to create CNVs (Step 2). Binding proteins (Step 1B). These DSBs are subsequently resolved by replication fork collapse (Step 1A) or cleavage by hairpin-formation relationship in intergenic regions (Table 3 and Figure S4). We detected elevated numbers of long A/T tracks across the genome. We hypothesize that one such benefit includes increased CNV generation and thus, increased adaptability especially in the face of antimalarial selection. In support of this hypothesis, the presence of CNV trigger sites across the genome poises every potential drug target for amplification (Figures 3 and 5). It is interesting to speculate that characteristics of CNV trigger sites could contribute to the observation that some clones develop resistance in vitro more readily than others (8,79). This would be the first time that DNA sequence itself, as opposed to the regulation of specific repair proteins (62,80), has been implicated in the ability of P. falciparum to develop resistance.

Potential DNA repair mechanisms leading to CNV formation in Plasmodium falciparum

Through the analysis of post-CN V sequences, we detect evidence for two DNA repair pathways acting in the generation of P. falciparum CNVs: microhomology-mediated end joining (MMEJ, (81-83)) and microhomology-mediated break-induced repair (MMBIR, (74,84)). The ubiquitous shortening of long A/T tracks after CNV generation as well as several single nucleotide insertions after repair implicates MMEJ, which can cause deletions with and without small insertions (clones D and F, Figure 4C, Supplementary Figure S4 and Supplementary Table S2). Alternatively, the presence of short repeat expansions points to MMBIR, which has not been characterized in P. falciparum (clones C and E, Figure 4D). Nucleotide addition is a common consequence of fork slippage during replication-mediated repair processes (74,82,84,85). Fork slippage is also a hallmark of an alternate and possibly unique pathway to P. falciparum, sequence of fork slippage during replication-mediated repair processes (86,87). Furthermore, when excluding homologous recombination as the distance of DNA resection which is the distance from DNA lesion to homologous sequence used for repair. For example, short-range resection biases repair toward microhomology-mediated pathways and extensive resection biases repair toward homologous recombination (86,87). Furthermore, when excluding homologous recombination, short resection distances of <50 bp are more likely to lead to MMEJ as a means of repair and longer distances <250 bp are more likely enacted by MMBIR (88). Our CNV ‘trigger site’ model suggests an important role for the A/T track-hairpin distance (Figure 5); we speculate that the span of sequence between each component could reflect the resection distance for either of these two repair pathways. Given the proposed 400 bp distance limit (Figure 3), there are 9130 intergenic and 2222 genic trigger sites capable of being utilized by these pathways (Table 3 and Figure 3). Although our study only assessed amplifications, repair of DSB breaks at these sites can lead to deletions as well but further investigation is required to understand the role of these pathways in CNV formation.

replication fork collapse (Step 1A) or cleavage by hairpin-binding proteins (Step 1B). These DSBs are subsequently repaired in a non-faithful manner to create CNVs (Step 2). Resulting amplifications are initially rare throughout P.falciparum populations but then undergo selection to remove deleterious CNVs and promote the maintenance of beneficial CNVs (Step 3).

CNV trigger sites are enriched within intergenic regions

We detected elevated numbers of long A/T tracks (>9 bp) and stable hairpins (>−5.8 kCal/mol) in intergenic regions when compared to genic regions of the P. falciparum genome (Table 2). Furthermore, we identified a closer track-hairpin relationship in intergenic regions (Table 3 and Figure 4) and a corresponding enrichment in trigger sites (defined as A/T tracks between 20 and 40 bp in length within 400 bp of a stable hairpin, which occurs for 19.0% of intergenic A/T tracks). These data indicate that there may be a selective benefit of their association in non-coding regions of the genome. We hypothesize that one such benefit includes increased CNV generation and thus, increased adaptability especially in the face of antimalarial selection. In support of this hypothesis, the presence of CNV trigger sites across the genome poises every potential drug target for amplification (Figures 3 and 5). It is interesting to speculate that characteristics of CNV trigger sites could contribute to the observation that some clones develop resistance in vitro more readily than others (8,79). This would be the first time that DNA sequence itself, as opposed to the regulation of specific repair proteins (62,80), has been implicated in the ability of P. falciparum to develop resistance.
Figure 5. Model of CNV development and selection in Plasmodium falciparum. In Step 1, DNA hairpins trigger double strand breaks throughout the P. falciparum genome presumably by either halting replication fork progression (Step 1A) or recognition by hairpin-binding proteins (Step 1B). In Step 2, long A/T tracks (gray circles) within 400 bp of the double strand break are utilized as microhomology for error-prone repair pathways to generate CNVs (blue, red and green bars). CNV breakpoints (vertical dotted lines) are generated semi-randomly across the genome but more stable hairpins are more likely to generate recurrent breakpoints (purple dotted line). De novo CNVs can either contain beneficial genes (gray bar with star) or those unrelated to the selection. New CNVs are generated frequently and could randomly occur throughout the highly repetitive P. falciparum genome (green bar), but may increase under selective pressure (see Discussion). In Step 3, selection (i.e. drug or fitness effects) enriches for beneficial CNVs (blue and red parasites) and purges deleterious CNVs (green parasite) from the population.

understand the mechanisms involved in the generation of deletions as well as how they contribute to the adaptability of the parasite.

Homologous recombination is highly active in the parasite (16,22,82,83); what then leads to the use of these error-prone pathways for repair? We propose that antimalarial treatment, which causes metabolic stress, skews repair toward MMEJ and MMBIR in P. falciparum. Microhomology-mediated pathways in other organisms have been shown to exhibit increased activity when cells are under stress (89–91). For example, under normal conditions in mammalian cells, RAD51 inhibits MMBIR activity and facilitates the use of homologous recombination for DSB repair (92). However, RAD51 is downregulated during hypoxic stress in tumors, dNTP depletion, and the starvation response in Escherichia coli and cancer as well as during replication stress in humans (90,92–96). Future studies on the levels of key repair proteins will be required to see if this is the case in P. falciparum.

Overall, we propose that a close A/T track-hairpin relationship in the P. falciparum genome leads to the utilization of error-prone microhomology-mediated pathways. These events lead to enhanced generation of CNVs and adaptability of this parasite under selective pressure. Further investigation of these mechanisms may identify DNA repair pathways that can be targeted to limit parasite adaptability.

SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online.

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