

# Genetic Evidence of Focal *Plasmodium falciparum* Transmission in a Pre-elimination Setting in Southern Province, Zambia

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**Background.** Southern Province, Zambia has experienced a dramatic decline in *Plasmodium falciparum* malaria transmission in the past decade and is targeted for elimination. Zambia's National Malaria Elimination Program recommends reactive case detection (RCD) within 140 m of index households to enhance surveillance and eliminate remaining transmission foci.

**Methods.** To evaluate whether RCD captures local transmission, we genotyped 26 microsatellites from 106 samples collected from index (n = 27) and secondary (n = 79) cases detected through RCD in the Macha Hospital catchment area between January 2015 and April 2016.

**Results.** Participants from the same RCD event harbored more genetically related parasites than those from different RCD events, suggesting that RCD captures, at least in part, infections related through local transmission. Related parasites clustered in space and time, up to at least 250 m from index households. Spatial analysis identified a putative focal transmission hotspot.

**Conclusions.** The current RCD strategy detects focal transmission events, although programmatic guidelines to screen within 140 m of index households may fail to capture all secondary cases. This study highlights the utility of parasite genetic data in assessing programmatic interventions, and similar approaches may be useful to malaria elimination programs seeking to tailor intervention strategies to the underlying transmission epidemiology.

**Keywords.** reactive case detection; malaria; microsatellites; screen-and-treat; molecular epidemiology.

The global decline in malaria between 2000 and 2015 has reinvigorated commitment to elimination [1]. Many countries have malaria elimination targets before 2030, including 8 in southern Africa. Although interventions such as long-lasting insecticide-treated bednets (LLINs), indoor residual spraying, and case management have been and remain important malaria control tools [1], transitioning from control to elimination will necessitate incorporating new strategies capable of permanently interrupting remaining avenues for local transmission [2].

Current elimination strategies recommend variations of active case detection (ACD) in which community health

workers (CHWs) screen community members for malaria using rapid diagnostic tests (RDTs) to identify asymptotically infected individuals who may contribute to ongoing transmission [3, 4]. Despite consensus that reducing the asymptomatic reservoir is theoretically important to elimination by preventing reestablishment of local transmission, the empirical evidence that screen-and-treat initiatives reduce the burden of malaria in practice is inconsistent [5–8]. While simulation analyses and a study from Zambia reported decreases in malaria infection following screen-and-treat efforts [5, 9], 2 cluster-randomized trials in Kenya and Burkina Faso found no difference in the change in malaria burden between villages receiving longitudinal screen-and-treat and villages receiving no intervention [7, 8].

Beyond conflicting evidence of effectiveness, ACD is costly and resource-intensive as an elimination strategy, particularly in low-transmission regions where few individuals have levels of parasitemia above the RDT limit of detection [10]. Therefore, reactive case detection (RCD), a specific type of ACD, has been adopted by numerous control programs in low-transmission settings, focusing screening efforts to geographic transmission “hotspots” rather than the entire community [3]. RCD, initiated

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after passive detection of an index case at a health facility, involves screening and treating individuals within the index household and in households a prescribed distance from the index household [3, 4]. Multiple studies have reported geographic clustering of malaria cases in such hotspots and noted that individuals living in index case households have a higher likelihood of being infected, supporting RCD as a strategy to disrupt transmission in ecological niches that support malaria transmission [11–19].

In Zambia, RCD was incorporated into the National Malaria Elimination Program (NMEP) in 2011 for regions reporting <1% RDT-positive prevalence, including Southern Province [20, 21]. In 2014, a ground-truth evaluation of RCD in Southern Province revealed that across 26 RCD events, only 32% of the households screened were actually located within the targeted 140-m radius, highlighting the challenges CHWs face in identifying eligible households [22]. Although RCD is theoretically appealing, the inconsistent data regarding ACD's ability to decrease community-level transmission coupled with the logistical challenges of RCD necessitates validating that RCD is appropriate for the malaria epidemiology of a given region. In particular, it is essential to determine whether clusters of malaria cases are sustained through local transmission or other mechanisms. Whereas local transmission may be effectively targeted through interventions like RCD and focal vector control, eliminating case clusters that arose from household travel or occupational exposure would require different approaches. Distinguishing these scenarios empowers evidence-based prioritization of appropriate interventions tailored to the underlying transmission epidemiology.

Spatial and temporal patterns of malaria cases, sometimes in conjunction with travel data, have been commonly used to draw inferences regarding the epidemiology of local malaria transmission [23–25]. Although they are an important aspect of epidemiological monitoring, spatial and temporal patterns, even when coupled with travel data, cannot reliably distinguish local transmission chains from clusters of unrelated infections. Understanding the contribution of local transmission requires comparing parasite genotypes from infected individuals [26, 27]. To evaluate the ability of RCD to capture local malaria transmission in a region of Southern Province, Zambia, we performed microsatellite genotyping on dried blood spot (DBS) samples collected from index and secondary cases detected through RCD. We characterized the temporal and spatial scales on which genetically related parasites clustered, a proxy for local transmission, and identified a putative focal transmission hotspot.

## METHODS

### Study Region

This study was conducted in the rural catchment area of Macha Hospital, an approximately 2000-km<sup>2</sup> region located in Choma District, Southern Province, Zambia, with an estimated 56 000 residents, primarily subsistence farmers. This setting is characterized

by a tropical savannah climate and experiences peak *Plasmodium falciparum* malaria during the single rainy season between November and April [28]. *Anopheles arabiensis* and *Anopheles squamosus* are the primary and secondary malaria vectors in this region, respectively [28–30]. *P. falciparum* histidine-rich protein 2 (PfHRP2) based RDT prevalence among actively detected individuals in this region declined from 9.2% in 2008 to <1% in 2013 [31]. The introduction of artemisinin combination therapy with artemether-lumefantrine in 2004 along with LLIN distributions in 2007, 2012, and 2014 may have contributed to the decline in malaria in this setting. LLIN ownership in this region was estimated to be 83% in 2013 [32]. The Zambian NMEP aims to eliminate malaria from the country by 2021, and Southern Province, including the study area, is considered a preelimination setting. For regions like this with low-level malaria transmission, the NMEP recommends RCD whereby CHWs screen using PfHRP2-based RDTs and treat positive individuals living within 140 m of index case households.

### Informed Consent

This research was approved by the Institutional Review Board at the Johns Hopkins Bloomberg School of Public Health (Baltimore, Maryland) and the Ethics Review Committee of the Tropical Diseases Research Centre (Ndola, Zambia). All adult participants gave informed consent. All children had parental consent, and children aged 12–15 provided assent.

### Enhanced Reactive Case Detection

Following passive detection of PfHRP2-RDT-positive index cases at 14 rural health centers (RHCs) serving the catchment area of Macha Hospital, study team members at Macha Research Trust (MRT) were alerted by text message. Within 1 week of the index case's presentation at the RHC, study team members accompanied a CHW to screen all individuals living within an expanded RCD radius of 250 m from the index case household using RDTs. RDT-positive cases were offered treatment with artemether-lumefantrine. For all individuals screened, study team members collected a finger-prick blood sample as DBS on filter paper (Whatman 903 Protein Saver Card), administered an epidemiological and demographic survey, and collected global positioning system coordinates for each household. Households within 250 m of the index household were preidentified using satellite imagery to guide the field team in enrolling eligible households. DNA from the DBS samples was extracted using Chelex and screened for *P. falciparum* using quantitative polymerase chain reaction (qPCR) targeting mitochondrial cytochrome-b at the MRT laboratory in Macha, Zambia [31].

### Passively Detected Cases at RHCs

Five of the 14 RHCs closest to Macha Hospital collected finger-prick blood samples as DBS from a convenience sample of passively detected index cases as previously described [33].

DBS samples were transported from the RHCs to MRT. Index cases were linked by name and visit date recorded at the RHC to the epidemiological and demographic survey questionnaire administered to the same individual up to 1 week later during RCD follow-up.

### Sample Selection

We selected all RCD events between 1 January 2015 and 18 April 2016 for which at least 2 participants, including the index case, were malaria positive by RDT or qPCR. One hundred six samples (27 index, 79 secondary) across 49 RCD events met our inclusion criteria, and DBS samples were transported to the University of California, San Francisco.

### Laboratory Procedures

DBS samples were stored with desiccant at  $-20^{\circ}$  C until transport and processing. Six-millimeter hole-punches from filter cards were extracted using the saponin Chelex method [34] and quantified using *var* gene acidic terminal sequence (*var*-ATS) ultra-sensitive qPCR [35]. Samples containing  $<10$  parasites/ $\mu$ L of blood (45/106 samples) underwent selective whole genome amplification (sWGA) to selectively enrich for parasite DNA as previously described, with some modifications (see [Supplementary Table 1](#)) [36, 37]. For quality control, we assessed the concordance of alleles and determined the error rate introduced during sWGA by genotyping control samples with and without sWGA. Samples with  $>10$  parasites/ $\mu$ L and post-sWGA samples were amplified at 26 microsatellite loci [38, 39] (Liu et al, unpublished data). PCR products were diluted and sized by denaturing capillary electrophoresis on a Thermo Fisher 3730XL analyzer with GeneScan 400HD ROX size standard (Thermo Fisher Scientific). The resulting electropherograms were analyzed using microSPAT (Murphy et al, unpublished data). Samples with  $<10$  successfully genotyped microsatellites and loci that did not amplify in at least 50% of the samples were excluded from the analysis. For each genotyped sample, we calculated the multiplicity of infection (MOI), or the number of genetically distinct parasite clones, as the second-highest number of alleles present at any locus, accounting for the possibility of false positive allele calls.

### Characterizing Relatedness Captured During RCD

For each pair of samples genotyped, pairwise genetic relatedness was determined using a modified identity by state (IBS) metric [40] (Liu et al, unpublished data). Pairwise IBS was calculated from allele similarity between isolates, allowing estimation of genetic relatedness from monoclonal as well as polyclonal samples. For each locus, IBS was estimated by the sum of the product of within host allele frequency (defined as  $1 / \text{number of alleles detected}$ ) of shared alleles. The overall pairwise IBS was determined as the average of locus-specific estimates under the assumption of independent loci (Liu et al, unpublished data).

We calculated the physical distance between all sample pairs as Euclidean distance between households using the R package “geosphere” [41]. Elapsed time between sample pairs was determined from the absolute value of the difference in sample collection dates. Sample pairs were considered related if their pairwise genetic relatedness was  $\geq 0.5$ , the 96th percentile of pairwise relatedness across all sample pairs. We characterized mean pairwise genetic relatedness as well as the proportion of related samples across bins of increasing physical distance. For each RCD event, we calculated the mean pairwise relatedness between all sample pairs from that event.

### Characterizing the Temporal and Spatial Boundaries of Malaria Transmission

We investigated the optimal temporal and spatial scales on which to detect related parasites in the study area. Given that the sampling framework inherently oversampled individuals clustered in time and space, we removed all pairwise comparisons between samples from the same RCD event. Using this subset of data corrected for the nonrandom sampling strategy, we binned the remaining pairwise comparisons into 9 space-time categories and calculated the mean genetic relatedness among pairwise comparisons across increasing time and distance.

### Identifying Focal Transmission Hotspots

We developed a method to identify regions that sustain local transmission at higher than expected rates, referred to as focal transmission hotspots. We used the R package “sp” [42–44] to draw lines between all pairs of individuals harboring related parasites after removing pairwise comparisons between samples from the same RCD event. We created a grid with 81 equally sized cells across the study region. For each cell, we counted the number of related lines that passed through, generating a count of observed related connections. To calculate an expected number of related connections for each cell under the assumption that connections were random, we generated 1000 sampling replicates of size equal to the total number of related lines, each time sampling from all possible lines connecting all pairs of individuals regardless of relatedness. For each replicate, we summed the number of lines that passed through each cell, generating an expected line count and 95% confidence interval for each cell. We calculated the ratio of observed to expected lines for each cell and exported this file for use in ArcMap (Esri, version 10.5.1) [45].

## RESULTS

### Genotyping

Of the 26 microsatellite loci amplified, all but locus PolyA had genotyping success rates  $>50\%$  ([Supplementary Figure 2](#)). Locus PolyA was therefore excluded from the analysis. Eighty-five of the 106 extracted samples were successfully genotyped for at least 10 of the 25 microsatellite loci and were included

in the analysis. Insufficient coverage was significantly associated with negative RDT result, being a secondary case, and low parasitemia (Supplementary Figure 3). Among the 21 samples with insufficient coverage, 19 underwent sWGA. The relationship between microsatellite coverage and parasitemia is shown in Supplementary Figure 3. Samples excluded from the analysis did not differ significantly from those included in terms of sex, age, self-reported overnight travel, or self-reported LLIN use. In a quality control experiment, there was >99% allele concordance in control samples genotyped pre- and post-sWGA (Supplementary Table 2).

The 85 successfully genotyped samples came from 47 different RCD events. Index ( $n = 27$ ) and secondary case individuals ( $n = 58$ ) did not differ significantly with respect to sex, age, self-reported LLIN use, or self-reported overnight travel (Supplementary Table 3). Index cases had a higher proportion of RDT-positive individuals and a lower proportion of sWGA samples compared to secondary cases, as expected given the higher parasite densities generally seen in symptomatic infections (Supplementary Table 3). Among the samples included, the median MOI was 2. MOI did not differ between index and secondary cases (Supplementary Figure 4) or between sWGA (median, 2 [range, 1–3]) and non-sWGA (median, 2 [range, 1–4]) samples.

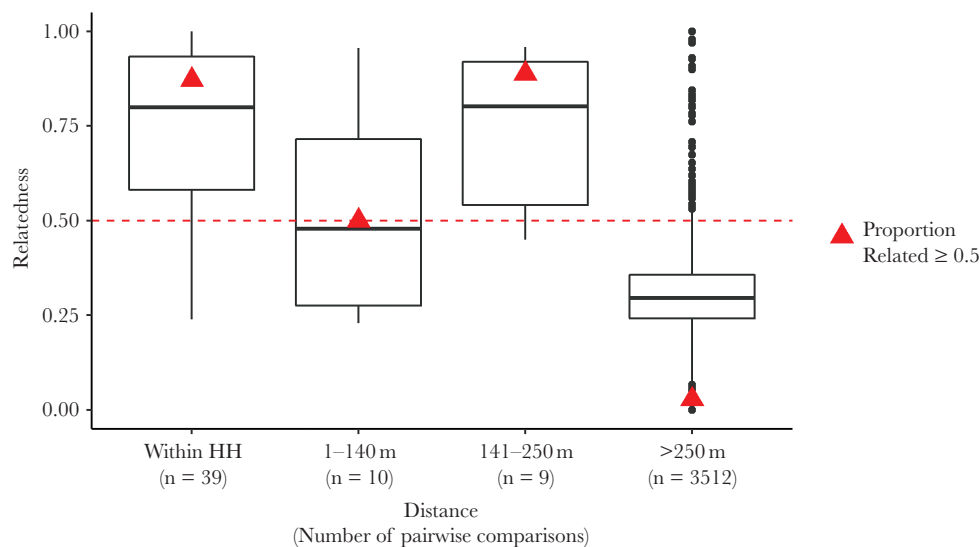
#### Relatedness by Physical Distance

There was no significant difference in mean genetic relatedness between samples collected from within the same household and those 141–250 m apart (Figure 1). Samples collected 1–140 m apart were significantly less related than those within the same household (Student  $t$  test,  $P = .004$ ) and those 141–250 m apart

(Student  $t$  test,  $P = .05$ ), but this difference may reflect the small sample size of 10 comparisons in the 1–140 m distance bin. When considered together, samples collected from participants located within 250 m had significantly higher mean genetic relatedness (mean relatedness, 0.71) than participants >250 m apart (mean relatedness, 0.31; Welch  $t$  test,  $P < .001$ ), indicating that screening with a 250-m radius captures genetically related parasites. Among samples from the same RCD event, 78.6% were related, whereas only 2.9% of samples from different RCD events were related (test for difference in proportions,  $P < .001$ ). The distributions of relatedness among sample pairs from the same RCD event and among those from different RCD events are shown in Supplementary Figure 1. Among the related pairwise comparisons >250 m apart, 60% occurred within 90 days, suggesting that relatedness among physically distant samples is nevertheless clustered temporally. In summary, RCD, both at the 140-m and 250-m radii, captured genetically related cases, suggesting that case clustering is due to local transmission rather than household-level risk factors.

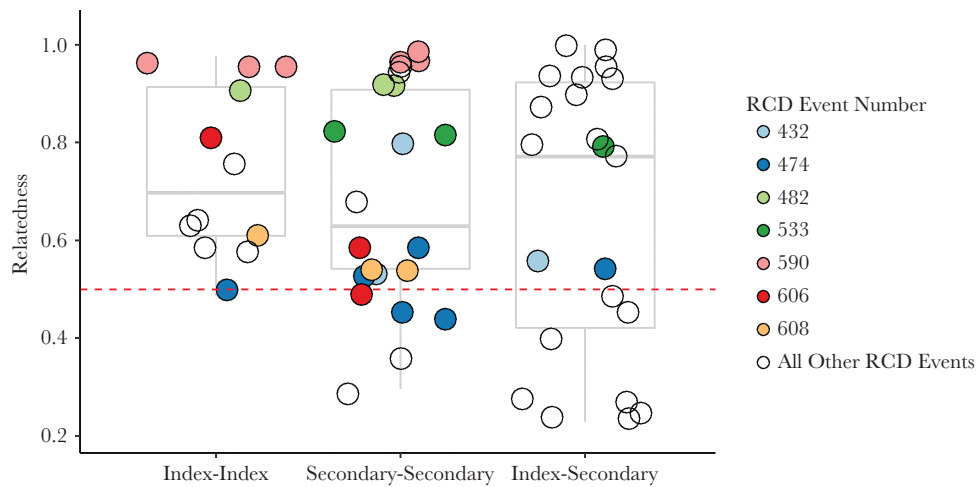
#### Relatedness Within RCD Events

Among the 56 sample pairs from the same RCD event, mean relatedness was 0.69. There was no significant difference in mean relatedness comparing 2 secondary cases, 2 index cases, or an index and secondary case from the same RCD event (analysis of variance,  $P = .62$ ; Figure 2). Among the 7 RCD events with comparisons across multiple categories (index–index, secondary–secondary, or index–secondary), there was no difference in relatedness across categories, possibly reflecting the small sample size.



**Figure 1.** Genetic relatedness declines after 250 m. Genetic relatedness is shown for pairwise comparisons binned by increasing physical distance between sample pairs (within the same household [HH], samples 1–140 m apart, samples 141–250 m apart, and samples >250 m apart). The number of pairwise comparisons in each physical distance bin is listed below the bin label in parenthesis. Triangles indicate the proportion of comparisons in each category that are related above the 0.5 genetic relatedness threshold, demarcated by the dashed line. Among sample pairs >250 m apart, the mean physical distance was 20.2 km (range, 297 m–46.1 km).



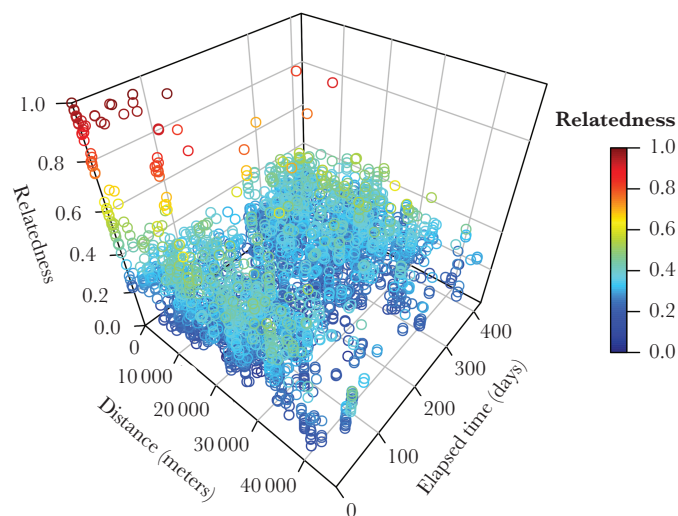


**Figure 2.** There were 56 pairwise comparisons between samples from the same RCD event, representing 23 unique RCD events. Each of the 56 comparisons were binned according to whether the comparison was between 2 index cases (index–index), between 2 secondary cases (secondary–secondary), or between an index case and a secondary case (index–secondary) from the same RCD event. Genetic relatedness is shown for each of the 56 comparisons, represented as circles. For 7 RCD events that had comparisons in >1 category (index–index, secondary–secondary, or index–secondary), comparisons are shaded identically. Pairwise comparisons from RCD events for which only 1 category is represented are shown with hollow circles. The distributions of all pairwise comparisons for each of the 3 categories are shown by boxplots.

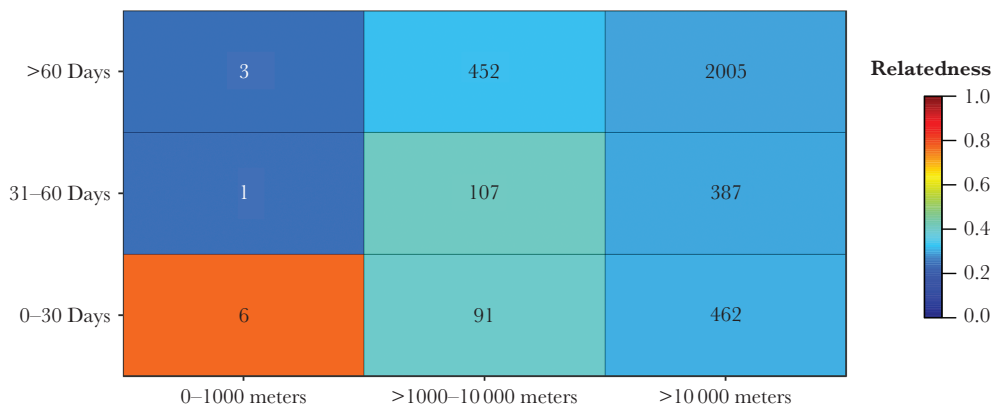
### Relatedness Across Space and Time

Related comparisons clustered focally in space and time (Figure 3), and pairwise genetic relatedness declined with increasing distance and elapsed time. Ninety-six percent of the 52 most highly related comparisons (related >0.80) were within 10 km and 92% occurred within 90 days (Figure 3), indicating that transmission occurred focally in space and time. Next, to verify that these relationships were not driven by the focal sampling framework, we removed all pairwise comparisons

between samples from the same RCD event. Figure 4 illustrates the decline in genetic relatedness across increasing distance and elapsed time after removing these comparisons. After excluding pairs of samples from the same RCD event, most related comparisons occurred within 10 km and 60 days, although the small sample size of comparisons within 1000 m precludes our ability to infer definitive boundaries between 0 and 1000 m. Marginal distributions of relatedness by physical distance and elapsed time are included in Supplementary Figure 5.



**Figure 3.** Relatedness declines across space and time. Each hollow circle represents a unique pairwise comparison between 2 samples. Circles are colored and positioned along the vertical axis according to the genetic relatedness between the samples compared. Relatedness of 1.0 (red) indicates that the samples are highly related, and relatedness of 0.0 (blue) indicates that the samples are unrelated. Physical distance and elapsed time between sample pairs are shown on the two horizontal axes.



**Figure 4.** Relatedness declines across space and time excluding same reactive case detection (RCD) event comparisons. Pairwise comparisons (excluding those from the same RCD event) were binned into 9 space–time categories represented by the 9 boxes in the matrix. The number in each bin denotes the number of pairwise comparisons in each category. The mean relatedness of the pairwise comparisons in each bin is shown by the color gradient, where relatedness of 1.0 (red) is highly related and relatedness of 0.0 (blue) is unrelated.

### Identifying Focal Transmission Hotspots

We identified a region in the northwest quadrant of our study area as a putative focal transmission hotspot (Figure 5). After removing pairs of samples from the same RCD event and accounting for the spatial distribution and density of households in the analysis, this region had a significantly higher than expected number of related connections. Conversely, 2 regions located in the west-central and north-central areas of our study site had significantly fewer than expected related connections. Parasite genetic data enabled the detection of finer-scale transmission heterogeneities in our study area than could have been detected through epidemiological and spatial patterns alone.

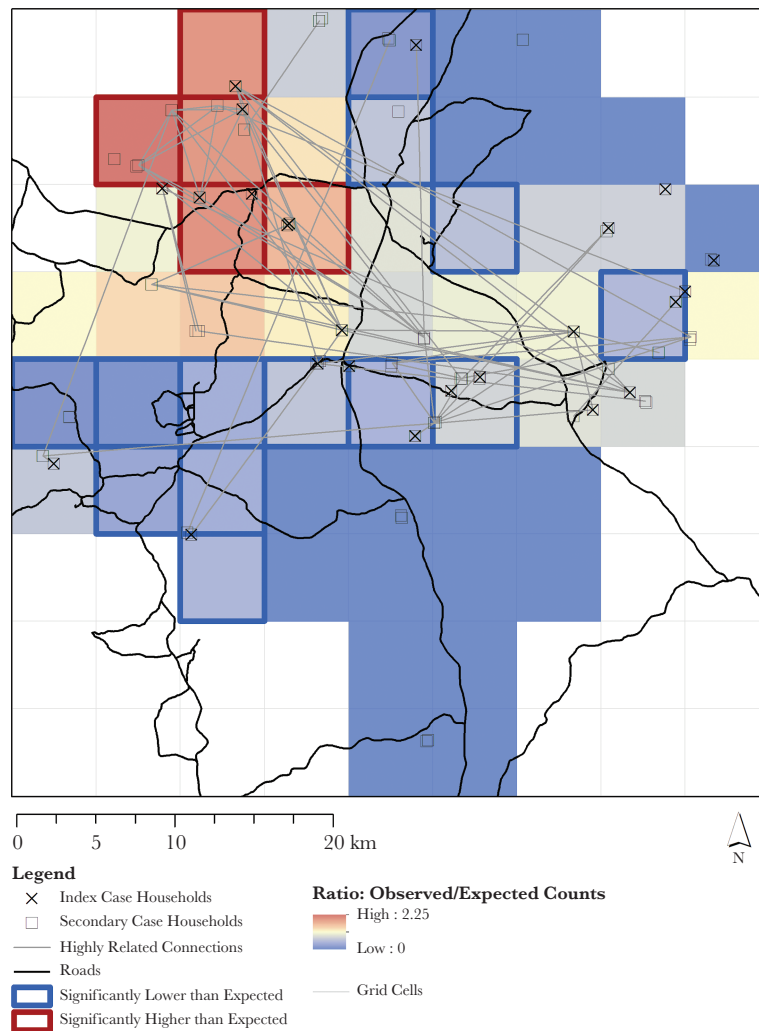
### DISCUSSION

Microsatellite genotyping from RCD participants demonstrated that focal malaria transmission was ongoing between January 2015 and April 2016 in a region of Southern Province, Zambia, targeted for elimination. Although spatial clustering of malaria cases has been previously described [11–14], this is, to our knowledge, the first study to genotype *P. falciparum* identified during RCD in southern Africa to attribute case clustering to local transmission. Using genetic methods to clarify the mechanism through which clusters of malaria cases arise has important implications for programmatic planning, enabling evidence-based decisions regarding effective control tools suited to a region’s unique malaria epidemiology. This study demonstrates that RCD captures genetically related infections and underscores the added value of genetic data to understanding the underlying malaria transmission epidemiology of a given region beyond what can be inferred through spatial and temporal patterns alone.

Our data, implicating local transmission as a contributor to the sustained malaria burden, highlights the importance

of addressing the underlying transmission biology and ecology that perpetuate malaria in this region. These results suggest that local transmission in this region extends up to 250 m from the index household, and possibly beyond. Therefore, the Zambian NMEP’s recommended 140-m screening radius at 1 timepoint likely fails to capture infected individuals who may be capable of sustaining transmission after RCD. However, performing RCD across the entire landscape of local transmission may not be feasible or efficient, as previously suggested [46, 47]. Exploring additional options, such as repeated visits or coupling focal vector control with RCD may be useful to interrupt local transmission in this and similar settings. With the goal of identifying regions where transmission-disrupting interventions such as focal vector control could have the largest impact, we describe a novel approach for detecting focal transmission hotspots. Focal transmission hotspots may be ideal locations to implement additional transmission-disrupting interventions in conjunction with RCD. This proposed method for identifying focal transmission hotspots has not yet been rigorously assessed and will require expanded evaluation before being considered a reliable indicator for transmission foci.

The RCD sampling framework in this study is not the ideal dataset to delineate the spatial and temporal scales of local transmission. We did not genotype every index and secondary case between January 2015 and April 2016, but rather selected samples that came from RCD events where at least 2 individuals tested positive for malaria by RDT or qPCR. Given the nonrandom inclusion criteria and somewhat sparse sampling of this study, the spatial and temporal boundaries of local transmission we can infer—10 km and 60–90 days—require additional validation. Despite our inability to unequivocally identify the spatial and temporal boundaries of local transmission, our



**Figure 5.** Observed vs expected counts of highly related connections. A map of the study area is overlaid with 81 equally sized, approximately 5-km grid cells. For each grid cell, the ratio comparing the number of related connections (pairwise comparisons related  $\geq 0.5$  after excluding comparisons from the same reactive case detection event) to the number of connections expected is depicted by a color gradient. A ratio  $> 1$  (red) indicates that the cell has more related connections than expected; a ratio  $< 1$  (blue) indicates that the cell has fewer related connections than expected. We observed a similar multiplicity of infection (MOI) comparing individuals living within and outside of focal transmission hotspots (mean MOI, 1.62 among individuals living in focal transmission hotspots vs 1.92 among individuals living outside of focal transmission hotspots; Student *t* test,  $P = .28$ ).

data strongly suggest that local transmission occurs focally in space and time. Temporal and spatial clustering was observed regardless of whether sample pairs from the same RCD event were included or excluded, indicating that our sampling framework did not solely determine the results. Our observations are consistent with simulation analyses from the region that suggested a 500-m RCD radius would identify 77% of additional cases [47]. Similarly, our observation that transmission occurs temporally for up to 60–90 days is corroborated by 24-SNP (single-nucleotide polymorphism) molecular barcode data suggesting ongoing transmission by highly related subsequent infections within index households 30 and 90 days after the initial RCD intervention [48]. Defining exact boundaries of local

transmission would necessitate sampling longitudinally from a spatially representative framework.

We inferred local transmission from pairwise genetic relatedness. Although we are unable to confidently assert that any sample pairs harboring related parasites represented direct transmission from one to another, it is improbable that we would see genetic relatedness values  $> 0.5$  unless the individuals were connected in some manner through transmission. Our 0.5 threshold represented the 96th percentile of all pairwise relatedness values in the study, indicating how rare these related events are. Despite not having the granularity of connecting individual transmission events, the observed related infections likely indicate direct or indirect local transmission.

The ability to obtain genotypes from asymptomatic infections, the diverse, multiallelic loci used, and the use of a similarity metric capable of comparing infections with varying MOI contributed to the success of this study in a setting of low-density, polyclonal infections. However, there is room for further improvement of these methods. For example, we were unable to assign meaning to categories of genetic relatedness values <0.5, given the low precision of characterizing how relatedness across transmission generations decays for these markers in the context of polyclonal infections.

Characterizing the contribution of local transmission to sustained malaria is critical to programs transitioning from control to elimination. Knowledge of whether cases arise through travel, occupational exposure, or locally acquired transmission can guide intervention priorities. We discerned signatures of local malaria transmission using 106 DBS samples and a relatively low-tech genotyping method. Similar approaches could be valuable to other researchers and program leaders, particularly in resource-limited settings, interested in characterizing local transmission to guide malaria elimination.

#### Supplementary Data

Supplementary materials are available at *The Journal of Infectious Diseases* online. Consisting of data provided by the authors to benefit the reader, the posted materials are not copyedited and are the sole responsibility of the authors, so questions or comments should be addressed to the corresponding author.

#### Notes

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