

# Efficacy of Artemether-Lumefantrine and Dihydroartemisinin-Piperaquine for Treatment of Uncomplicated Malaria in Children in Zaire and Uíge Provinces, Angola

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**The development of resistance to antimalarials is a major challenge for global malaria control. Artemisinin-based combination therapies, the newest class of antimalarials, are used worldwide but there have been reports of artemisinin resistance in Southeast Asia. In February through May 2013, we conducted open-label, nonrandomized therapeutic efficacy studies of artemether-lumefantrine (AL) and dihydroartemisinin-piperaquine (DP) in Zaire and Uíge Provinces in northern Angola. The parasitological and clinical responses to treatment in children with uncomplicated *Plasmodium falciparum* mono-infection were measured over 28 days, and the main outcome was a PCR-corrected adequate clinical and parasitological response (ACPR) proportion on day 28. Parasites from treatment failures were analyzed for the presence of putative molecular markers of resistance to lumefantrine and artemisinins, including the recently identified mutations in the K13 propeller gene. In the 320 children finishing the study, 25 treatment failures were observed: 24 in the AL arms and 1 in the DP arm. The PCR-corrected ACPR proportions on day 28 for AL were 88% (95% confidence interval [CI], 78 to 95%) in Zaire and 97% (91 to 100%) in Uíge. For DP, the proportions were 100% (95 to 100%) in Zaire, and 100% (96 to 100%) in Uíge. None of the treatment failures had molecular evidence of artemisinin resistance. In contrast, 91% of AL late-treatment failures had markers associated with lumefantrine resistance on the day of failure. The absence of molecular markers for artemisinin resistance and the observed efficacies of both drug combinations suggest no evidence of artemisinin resistance in northern Angola. There is evidence of increased lumefantrine resistance in Zaire, which should continue to be monitored.**

Successful malaria control and elimination programs rely on the availability of effective antimalarials. However, resistance to antimalarials poses a threat to the global effort in malaria control. Widespread resistance to conventional antimalarials, such as chloroquine, sulfadoxine-pyrimethamine, and mefloquine, motivated the World Health Organization (WHO) to recommend the use of artemisinin-based combination therapy (ACT) for the treatment of uncomplicated *Plasmodium falciparum* malaria (1). Although ACTs were introduced only in the 1990s, studies have documented the emergence of resistance to artemisinins in limited areas of the Greater Mekong Subregion (2–10), and a recent dispatch reported a case of frank artemisinin treatment failure in a traveler from Angola (11).

Due to the risk of the emergence and spread of antimalarial drug resistance, the WHO recommends that countries where malaria is endemic routinely retest the efficacy of first-line antimalarials using a standardized protocol (12). Reduced efficacy of ACTs can be due to resistance to the fast-acting artemisinin derivative, the longer-lasting partner drug, or both. Resistance to the artemisinin derivative is typically characterized by delayed parasite clearance. In contrast, resistance to the partner drug more often manifests as recrudescence infections, occurring more than 7 days after the start of treatment, following initial parasite clearance. As a complement to clinical evidence, parasites from treatment failures can be tested for the presence of molecular markers of resistance to the component drugs. Resistance to the commonly used partner drug lumefantrine has been tentatively associated with certain specific mutations in the *pfmdr1* gene (13). For arte-

misinin resistance, a promising molecular marker in the propeller domain of the kelch protein, K13, was recently identified (9). Mutations in this gene have been found in resistant parasites in samples from Southeast Asia, the same region where resistance to chloroquine and sulfadoxine-pyrimethamine first arose and then spread to Africa. Mutations in this gene have also been found in a small proportion of samples from Africa, but none of these matched mutations associated with delayed clearance in Southeast Asia (10).

Angola, a country in Southern Africa where malaria is endemic, began recommending ACTs as first-line treatment for uncomplicated malaria in 2005. Currently, three ACTs are equally recommended for use in Angola: artemether-lumefantrine (AL),

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artesunate-amodiaquine (ASAQ), and dihydroartemisinin-piperazine (DP). The most recent data on the efficacy of ACTs in Angola comes from a 2004 *in vivo* therapeutic efficacy study that found 100% PCR-corrected efficacy of ASAQ and AL in Huambo Province in central Angola (14).

Here, we report the results of back-to-back *in vivo* studies of AL and DP carried out in both Zaire and Uíge Provinces in northern Angola in 2013 designed to measure the therapeutic efficacy of each drug for the treatment of uncomplicated *P. falciparum* malaria. In addition, we report the results of screening the treatment failures for the presence of molecular markers associated with lumefantrine and artemisinin resistance.

## MATERIALS AND METHODS

We followed the standard 28-day WHO protocol for *in vivo* therapeutic efficacy studies (12). The study was designed and powered to independently measure the efficacy of AL and DP in each province, for a total of four arms. The study was not designed to compare the two drugs; there was no randomization, and in each province patients were first enrolled in the AL arm until the target number was reached, and then patients were enrolled in the DP arm.

**Study location and population.** Malaria transmission in Zaire Province, on the Atlantic coast in the northwest corner of the country, is classified as mesoendemic stable, while in Uíge Province, directly to the west of Zaire, transmission is classified as hyperendemic. Both provinces border the Democratic Republic of the Congo to the north. Enrollment and follow-up of patients took place between February and May of 2013 and coincided with the peak malaria transmission season.

In Zaire Province, the study took place in the outpatient clinics of the M'Banza Congo Municipal Hospital and the 11 de Novembro Maternal and Child Health Center, both located in the provincial capital. In Uíge Province, participants were enrolled and followed in the outpatient clinic of the General Hospital of Uíge, the main referral hospital for the province. Participants were children between 6 and 108 months (9 years) of age and were restricted to those living in close proximity to the clinics (generally within 5 km). To achieve a precision of 5% assuming an expected efficacy of 95%, the minimal sample size was calculated to be 73 for each arm, and 100 participants were planned to be enrolled to allow for loss to follow-up.

**Study procedures.** Children in the specified age category coming into the outpatient clinics with fever or history of fever within 24 h were identified and screened for eligibility to participate in the study. Children were eligible to participate if they had uncomplicated *P. falciparum* mono-infection with a parasite density of between 2,000 and 100,000 asexual parasites/ $\mu$ L. Children were excluded from participating if they weighed less than 5 kg, were malnourished (defined as a weight for age Z-score of <3), were severely anemic (hemoglobin less than 5 g/dl), reported having taken drugs with antimalarial activity in the previous 2 weeks, had general danger signs or signs of severe malaria, or if their guardians were unwilling or unable to bring them back for all follow-up visits.

The first 100 children enrolled in each province were treated with AL (Coartem-D dispersible tablets; Novartis, Basel, Switzerland). The next 100 children were treated with DP (Duo-cotecxin; Beijing Holley-Cotec, Beijing, China). The manufacturers' weight-based dosing schedules were followed. If the child vomited within a half hour of taking the medication, the entire dose was repeated. If the child vomited between a half hour and an hour after taking the medication, a half-dose was given. Children who vomited more than once in 24 h were excluded. While all three doses of DP were directly observed in the clinic, three of the AL doses were observed in the clinic, and the three evening doses were given at home. The children's parents or guardians were given a dose to administer at night, together with a reserve dose in case of vomiting. Patients with cell phones were called twice in the evening during treatment days—once to remind them about giving the dose and 1 h later to inquire about vomiting.

Clinical and parasitological responses to treatment were assessed over

a 28-day follow-up, with clinic visits including a clinical exam on days 0, 1, 2, 3, 7, 14, 21, and 28. The drugs were administered on days 0, 1, and 2. Parasitemia was measured through microscopy on days 0, 2, 3, 7, 14, 21, and 28 of follow-up. Slides were read by two technicians posted in each of the three participating health facilities, following standard WHO procedures (12). A subset of slides was quality controlled during periodic supervisory visits by an external reader. Hemoglobin was measured on days 0, 14, and 28 using a HemoCue (AB Leo Diagnostics, Helsingborg, Sweden) machine. Dried blood spots were collected on days 0, 7, 14, 21, and 28 for potential molecular analysis and were stored in individual plastic bags with desiccant at room temperature for later transport to CDC laboratories in Atlanta, GA.

Participants were classified as early treatment failures if they developed signs of severe malaria in the presence of parasitemia up to day 3, if day 2 parasitemia was higher than day 0 parasitemia, if day 3 parasitemia was higher than 25% of day 0 parasitemia in the absence of fever, or if there was any parasitemia on day 3 in the presence of fever. Participants were classified as late treatment failures if they did not demonstrate early treatment failure and had at least 1 blood slide positive for asexual *P. falciparum* parasites after day 3.

Patients meeting criteria of treatment failure were treated with intravenous quinine and removed from further participation in the study. Patients negative for malaria on day 28 and not having previously met any treatment failure criteria were classified as having an adequate treatment response.

**Laboratory procedures.** Genomic DNA was isolated from dried blood spots using the QIAamp DNA blood minikit (Qiagen, Valencia, CA) per the manufacturer's instruction. Confirmation of *P. falciparum* infection and DNA quality assessment were conducted using photoinduced electron transfer (PET)-PCR (15).

A panel of seven neutral microsatellite markers on six chromosomes was chosen to determine the genotype on the parasite genetic background and compare the genotype profiles of parasites collected at the time of enrollment and at the time of failure (16–18). The fragments were amplified by PCR, separated on an ABI3130xl genetic analyzer, and scored using GeneMapper v3.1 (Applied Biosystems, Foster City, CA) according to previously described methods (19). An allele was considered different when the length of the allele was more than 2 bp different. Patient samples from day 0 and day of failure sharing allele sizes for at least six of seven neutral microsatellite loci were considered recrudescence infections (20, 21). The multiplicity of infection was defined as the maximum number of alleles at a single locus from a single sample.

Putative markers of lumefantrine and artemisinin resistance were also examined for all treatment failures on day 0 and day of failure (no known molecular markers for piperazine exist). The *pfmdr1* copy number was estimated by real-time PCR following a previously described method (22). Two regions of *pfmdr1*, covering codons 86 to 184 and 1034 to 1246, were amplified and analyzed for polymorphisms by direct sequencing using a previously described method (23). The *pfmdr1* haplotypes were constructed for treatment failures from the *pfmdr1* sequences. Previously reported polymorphisms associated with artemisinin resistance on chromosome 10 (MAL10-688956) and chromosome 13 (MAL13-1718319) (8) were also assayed by PET-PCR and pyrosequencing. In addition, the recently identified artemisinin resistance markers on the kelch K13-propeller gene were assayed using Sanger sequencing (see the methods in the supplemental material).

**Statistical and sequence analyses.** Double data entry was done daily using Epi Info 7 (CDC, Atlanta, Georgia). All statistical analysis was done using R version 2.15.1 (R Foundation for Statistical Computing, Vienna, Austria). All sequence analysis was done using Geneious Pro R7 (Biomatters, Inc., Auckland, New Zealand).

The main outcome analyzed was the 28-day PCR-corrected proportion of adequate clinical and parasitological response (ACPR), defined as the fraction of participants that finished the study that were classified as having adequate treatment responses, excluding any patients with PCR-

**TABLE 1** Characteristics of patients at enrollment in therapeutic efficacy studies in Zaire and Uíge Provinces, Angola, in 2013

Parameter	<i>n</i> (%) in study arm			
	AL		DP	
	Zaire	Uíge	Zaire	Uíge
Initial screen	305	188	329	230
Slide read	287 (94)	173 (92)	326 (99)	222 (97)
Parasitemia (parasites/ml)				
Negative for plasmodium	112 (39)	53 (31)	174 (53)	78 (35)
Low (<2,000)	23 (8.0)	10 (5.8)	20 (6.1)	15 (6.7)
Target parasitemia (2,000–100,000)	102 (36)	101 (58)	104 (32)	102 (46)
High parasitemia (>100,000)	50 (17)	9 (5.2)	28 (8.6)	28 (13)

determined reinfection or any patients excluded over the course of the study. Secondary outcomes included (i) PCR-uncorrected 28-day ACPR, where cases of reinfection were included as treatment failures, and (ii) parasite clearance rates for day 3, defined as the proportion of evaluable patients without any asexual parasites on that day.

Kaplan-Meier estimates of the survival function for each arm were calculated, and *P* values corresponding to the significance of differences in time to failure between provinces were calculated from the log rank test. For this analysis, data from patients lost to follow-up or patients excluded from the per-protocol analysis due to protocol violation were included up until the final visit before exclusion and censure. In the PCR-corrected analysis, cases of reinfections were censored at the day of reinfection. The PCR-corrected and PCR-uncorrected cumulative success rates on day 28 were estimated using the Kaplan-Meier estimator, with confidence intervals calculated using the Greenwood method.

**Ethical considerations.** Official human subject oversight approval to conduct the evaluation was obtained from the Centers for Disease Control and Prevention in Atlanta, GA, and the National Malaria Control Program in Angola. Written informed consent was obtained from participants' parents or guardians before enrollment.

## RESULTS

**Screening.** The target enrollment numbers of 100 children in each arm were reached within 2 months in both provinces. Malaria positivity in screened children ranged from 47% to 69% in the four arms (Table 1). At least 5% of tested children had a high parasitemia (>100,000 parasites/ $\mu$ l) in each arm. In order to enroll 100 children for each arm, we needed to screen between 188 and 329 children for each arm. The most common cause of exclusion during screening was the presence of no parasitemia.

**Outcomes.** The rates of loss to follow-up were 11% and 10% in the AL arms in Zaire and Uíge, respectively, and 20% and 9% in the DP arms in Zaire and Uíge, respectively (Table 2). There were a total of 32 patients excluded from the study after enrollment. Reasons for this included the incorrect dose taken or dose not

taken at home (12), persistent vomiting (8), taking another anti-malarial after enrollment (7), incorrect enrollment of a child not meeting the inclusion criteria (3), and development of signs of severe illness in the absence of parasitemia (2).

This resulted in 79 and 78 children finishing the study in the AL arms in Zaire and Uíge, respectively, and 80 children in the DP Zaire arm and 83 children in the DP Uíge arm (Table 3). There were 25 treatment failures documented during the study: 24 in the AL arms and 1 in the DP arms. One of the treatment failures was an early treatment failure from the AL Zaire arm; in this case the child's hemoglobin (initially 5.5 g/dl) dropped below 5 g/dl on day 2, with a parasitemia of 615 parasites/ $\mu$ l. The PCR-uncorrected ACPR rates were 77.2% (95% confidence interval [CI], 66 to 86%) in the AL Zaire arm, 92.3% (84 to 97%) in the AL Uíge arm, 98.8% (93 to 100%) in the DP Zaire arm, and 100% (96 to 100%) in the DP Uíge arm. Time to failure was significantly shorter in the AL Zaire arm than in the AL Uíge arm in the PCR-uncorrected analysis, with a *P* value of 0.008 (Fig. 1). The PCR-uncorrected cumulative success rates on day 28 for each study arm closely matched the ACPR proportion on day 28 (Table 3).

Using microsatellite genotyping analysis (see Table S1 in the supplemental material), we were able to classify 15 of the 24 late treatment failures as reinfections and 9 as recrudescence infections, with 7 of the recrudescence infections occurring in the AL Zaire arm and the remaining 2 in the AL Uíge arm. This translated to 28-day PCR-corrected ACPR proportions of 88.4% (95% CI, 78 to 95%) in the AL Zaire arm, 97.3% (91 to 100%) in the AL Uíge arm, 100% (95 to 100%) in the DP Zaire arm, and 100% (96 to 100%) in the DP Uíge arm. The survival curves for the AL arms in the two provinces were also significantly different in the PCR-corrected analysis, with a *P* value of 0.05 (Fig. 1). The PCR-corrected cumulative success rates on day 28 for each study arm closely matched the ACPR proportion on day 28 (Table 3).

For AL, day 3 clearance rates were 100% (95% CI, 96 to 100%) in Zaire and 97.6% (92 to 100%) in Uíge. For DP, all patients in both provinces cleared their parasitemia by day 3 (Table 3).

**Molecular analyses.** The 25 treatment failures were further characterized using previously identified molecular markers associated with drug resistance to either lumefantrine or artemisinin (Table 4). All failures had a single copy of *pfmdr1*. Sequencing of the *pfmdr1* gene only showed mutations at codons 86, 184, and 1246. Haplotypes constructed from these codons reflected haplotypes previously associated with lumefantrine resistance (13). While six different *pfmdr1* haplotypes were observed in day 0 samples, only three haplotypes were present in day of failure samples. The NFD haplotype was found

**TABLE 2** Sample size by study arm in therapeutic efficacy studies in Zaire and Uíge Provinces, Angola, in 2013

Parameter	<i>n</i> (%) in study arm			
	AL		DP	
	Zaire	Uíge	Zaire	Uíge
Total enrolled	101	99	102	100
Lost to follow-up	11 (11)	10 (10)	20 (20)	9 (9)
Excluded/censored	11 (11)	11 (11)	2 (2)	8 (8)
Finished study	79 (78)	78 (79)	80 (78)	83 (83)

**TABLE 3** Response to treatment in therapeutic efficacy studies in Zaire and Uíge Provinces, Angola, in 2013

Parameter	Result in study arm			
	AL		DP	
	Zaire ( <i>n</i> = 79)	Uíge ( <i>n</i> = 78)	Zaire ( <i>n</i> = 80)	Uíge ( <i>n</i> = 83)
ACPR, <i>n</i> (%)	61 (77)	72 (92)	79 (99)	83 (100)
Treatment failure, <i>n</i> (%)	18 (23)	6 (7.7)	1 (1.2)	0 (0)
Early	1 (1.3)	0 (0)	0 (0)	0 (0)
Late	17 (22)	6 (7.7)	1 (1.2)	0 (0)
Day 14	1 (1.3)	0 (0)	0 (0)	0 (0)
Day 21	10 (13)	3 (3.8)	0 (0)	0 (0)
Day 28	6 (7.6)	3 (3.8)	1 (1.2)	0 (0)
Reinfections	10 (13)	4 (5.1)	1 (1.2)	0 (0)
Recrudescence	7 (8.9)	2 (2.6)	0 (0)	0 (0)
Day 3 clearance, % (95% CI)	100 (96–100)	97.6 (92–100)	100 (96–100)	100 (96–100)
Cumulative success rate on day 28, % (95% CI)				
PCR uncorrected	77.4 (69–87)	92.3 (87–98)	98.8 (96–100)	100 <sup>a</sup>
PCR corrected	89.6 (83–97)	97.4 (94–100)	100 <sup>a</sup>	100 <sup>a</sup>
Proportion of ACPR on day 28, % (95% CI)				
PCR uncorrected	77.2 (66–86)	92.3 (84–97)	98.8 (93–100)	100 (96–100)
PCR corrected	88.4 (78–95)	97.3 (91–100)	100 (95–100)	100 (96–100)

<sup>a</sup> Confidence intervals were not calculated for the Kaplan-Meier estimator when the rate was 100%.

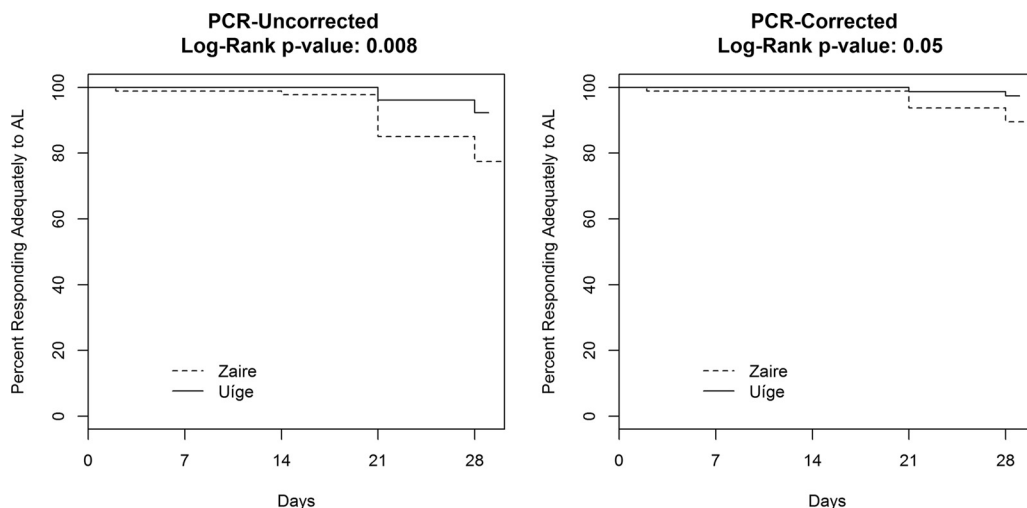
on day of failure in four recrudescences (44% of AL recrudescences) and four reinfections (31% of AL reinfections). The NYD haplotype was found on day of failure in five recrudescences (56%) and eight reinfections (54%). The YYD haplotype was only found on the day of failure in two reinfections (15%).

All isolates, both on day 0 and the day of treatment failure, were wild type at loci on chromosomes 10 and 13, where single-nucleotide polymorphisms (SNPs) had originally been shown to be associated with artemisinin resistance. The kelch K13 propeller gene, the site of the recently identified molecular markers of artemisinin resistance, was wild type for all day 0 and day of treatment failure isolates.

**Conclusions.** The 28-day PCR-corrected efficacies for the AL

Uíge arm and both DP arms are consistent with previously reported data for AL (24, 25) and DP (26, 27) and with high clinical efficacy estimates measured in African children prior to the widespread use of ACT. The point estimate for the 28-day PCR-corrected ACPR proportion in the AL Zaire arm of 88.4% is below the 90% WHO target efficacy for first-line treatment policy (12), but the 95% confidence interval of 78 to 95% is wide and includes the 95% secondary WHO target efficacy for ACTs. While the efficacies of DP are uniformly high in both provinces, AL is significantly less efficacious in Zaire Province than in Uíge Province.

The true efficacy of AL could be underestimated here, as we were not able to observe the three evening doses. However, previous studies comparing the efficacy of fully supervised and unsu-



**FIG 1** PCR-uncorrected and PCR-corrected survival functions for time until failure from a 2013 therapeutic efficacy study of AL in Zaire and Uíge Provinces, Angola. DP data are not shown as efficacy was uniformly high.

TABLE 4 Molecular characteristics of 25 observed treatment failures during therapeutic efficacy studies in Zaire and Uíge Provinces, Angola, in 2013

Patient ID no.	Treatment arm <sup>a</sup>	Classification <sup>b</sup>	MOI <sup>c</sup>	Resistance marker(s) on day 0 and day of failure <sup>d</sup>				
				<i>pfmdr1</i> <sup>e</sup>		SNP <sup>f</sup>		
				CN	Haplotype	Chr 10	Chr 13	K13
B313 <sup>g</sup>	AL-Z	ETF	2	1	NYD	wt	wt	wt
A111	AL-U	RECR	1	1	NFD	wt	wt	wt
A145	AL-U	RECR	1	1	NYD	wt	wt	wt
B314	AL-Z	RECR	1	1	NYD	wt	wt	wt
B384	AL-Z	RECR	1	1	NYD	wt	wt	wt
B385	AL-Z	RECR	2	1	NFD	wt	wt	wt
B399	AL-Z	RECR	2	1	NFD	wt	wt	wt
B404	AL-Z	RECR	2	1	NFD	wt	wt	wt
B416	AL-Z	RECR	2	1	NYD	wt	wt	wt
B422	AL-Z	RECR	2/1	1	NYD+NFD/NYD	wt	wt	wt
A114	AL-U	REIN	1	1	NYD	wt	wt	wt
A115	AL-U	REIN	2/1	1	YFD/YYD	wt	wt	wt
A144	AL-U	REIN	4/1	1	NYD	wt	wt	wt
A171	AL-U	REIN	1/2	1	YYD/NFD	wt	wt	wt
B304	AL-Z	REIN	1	1	NFD	wt	wt	wt
B312	AL-Z	REIN	1	1	NYD	wt	wt	wt
B371	AL-Z	REIN	2	1	NFD	wt	wt	wt
B375	AL-Z	REIN	1/2	1	NFD/YYD	wt	wt	wt
B386	AL-Z	REIN	1	1	YYY/NYD	wt	wt	wt
B387	AL-Z	REIN	1	1	NFY/NYD	wt	wt	wt
B398	AL-Z	REIN	1	1	NYD	wt	wt	wt
B402	AL-Z	REIN	2/1	1	YYY/NYD	wt	wt	wt
B421	AL-Z	REIN	1/2	1	NYD/NFD	wt	wt	wt
B423	AL-Z	REIN	1/2	1	NFD/NYD	wt	wt	wt
C560	DP-Z	REIN	1	1	YYD	wt	wt	wt

<sup>a</sup> AL, artemether-lumefantrine; DP, dihydroartemisinin-piperazine; Z, Zaire; U, Uíge.

<sup>b</sup> ETF, early treatment failure; RECR, recrudescence; REIN, reinfection.

<sup>c</sup> Multiplicities of infection (MOI) either are the same for day 0 and the day of failure or are reported as day 0 MOI/day of failure MOI.

<sup>d</sup> Resistance markers either are the same for day 0 and the day of failure or are reported as day 0/day of failure. wt, wild type.

<sup>e</sup> "CN" represents the *P. falciparum* multidrug resistance gene 1 copy number, and "haplotype" represents the haplotype constructed from mutations N86Y, Y184F, and D1246Y.

<sup>f</sup> Shown are the chromosome 10 (Chr 10) SNP at position 688956 and Chr 13 SNP at position 1718319 associated with artemisinin resistance.

<sup>g</sup> Molecular data for early treatment failure are available for day 0 only.

pervised (five out of six doses taken at home) dosing did not find significant differences in measured efficacies (28, 29).

Meta-analyses show that DP has higher PCR-corrected efficacies than AL, at least in the African setting (30). Moreover, DP has consistently been shown to have a lower reinfection rate in 28-day follow-up studies (30), likely due to the difference in elimination half-lives between piperazine (23 days in adults) and lumefantrine (4.5 days in adults) (31, 32).

Because of nonrandom assignment to the AL and DP arms, we cannot statistically compare the efficacies of the two drugs in this setting. Participants were enrolled in the DP arms only after the AL enrollment was finished, and consequently the follow-up for the DP arms systematically occurred later in the transmission period. Therefore, the smaller number of reinfections in the DP arm might be due to a lower transmission intensity and a lower risk of reinfection. The 28-day follow-up period might also mean some DP treatment failures went unobserved because of the longer half-life of piperazine.

Our finding of a single *pfmdr1* copy number is similar to those from previous studies from Africa, where a single *pfmdr1* copy number has been found in isolates from patients who experienced AL treatment failure (13). We found high rates of two *pfmdr1* haplotypes containing mutations previously associated with lu-

mefantrine resistance, NFD and NYD, in isolates from reinfections and recrudescences from the AL arms in both provinces. Our results are consistent with data from Uganda showing selection of the 86N, 184F, and 1246D mutations following treatment with AL (33, 34). Malmberg et al. recently reported that parasites with the *pfmdr1* NFD and NYD haplotypes could tolerate 15- and 7-fold-higher lumefantrine blood concentrations, respectively, than parasites with the YYY haplotype (35). Overall, 100% of day of failure isolates from recrudescences and 87% of day of failure reinfections in the AL arms had either an NFD or NYD haplotype.

The *pfmdr1* results, together with the uniformly wild-type chromosome 10 and 13 and kelch K13 propeller data and parasite clearance rates, suggest that the observed failure rates of AL are likely due to lumefantrine resistance in circulating parasites. The significant difference in efficacy of AL between provinces suggests that rates of lumefantrine resistance might be higher in Zaire than Uíge.

The measured efficacy and parasite clearance rates of both drugs, combined with the confirmed absence of putative molecular markers of artemisinin resistance, are consistent with a parasite population still exquisitely sensitive to artemisinins. There is sufficient evidence for the continued use of AL and DP in northern Angola. Although the measured efficacy of AL in Zaire Province is

below the 90% threshold at which WHO suggests changing treatment policy, a single estimate with wide confidence intervals and limited in time and space does not necessarily provide sufficient evidence for policy change. However, the efficacy of AL and molecular markers of resistance should be monitored in Zaire Province and at other sites across the country due to the apparent existence of lumefantrine resistance in the parasite population and the potential for selective pressure to intensify it. A follow-up therapeutic efficacy study of AL is planned to take place in Zaire Province and other sites in early 2015. These data should help provide a more comprehensive picture of antimalarial resistance and inform decisions regarding national treatment policy in Angola.

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