Molecular identification of the *Anopheles nili* group of African malaria vectors

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**Abstract.** Distinction between members of the *Anopheles nili* group of mosquitoes (Diptera: Culicidae), including major malaria vectors in riverside villages of tropical Africa, has been based mainly on doubtful morphological characters. Sequence variations of the ribosomal DNA second internal transcribed spacer (ITS2) and D3 28S region between morphological forms revealed four genetic patterns corresponding to typical *An. nili* (Theobald), *An. carnewei* Brunhes et al., *An. somalicus* Rivola & Holstein and the newly identified variant provisionally named Oveng form. Primers were designed based on ITS2 fixed nucleotide differences between haplotypes to develop a multiplex PCR for rapid and specific identification of each species or molecular form. Specimens of the *An. nili* group from Cameroon, Burkina Faso, Ivory Coast and Senegal were successfully identified to species, demonstrating the general applicability of this technique based on criteria described in this paper.

**Key words.** *Anopheles nili*, allele-specific PCR, malaria vectors, ribosomal DNA, sibling species, Afrotropical, Cameroon, Burkina Faso, Ivory Coast, Senegal.

**Introduction**

*Anopheles* (*Cellia*) mosquitoes are responsible for human malaria transmission in tropical areas of the Old World. In Africa, most of the important malaria vectors are members of species complexes, whose members are difficult and sometimes impossible to distinguish morphologically (Gillies & De Meillon, 1968; Coluzzi, 1984; Gillies & Coetzee, 1987; Fontenille & Lochouarn, 1999). These difficulties have stimulated the development of molecular tools for precise and reliable identification of sibling species. One of the most widely used regions of the genome to infer genetic variation and phylogenetic relationships is the ribosomal DNA (rDNA) cluster, a tandemly repeated multigene family. In eukaryotic organisms, every repeated unit of rDNA consists of an intergenic spacer (IGS), followed by genes coding the 18S, 5.8S and 28S rDNA. Preceding the 18S gene is the external transcribed spacer (ETS), and surrounding the 5.8S rDNA are the internal transcribed spacers 1 and 2 (ITS1 and ITS2). This multigene family evolves cohesively within species through concerted evolution, a mechanism that tends to homogenize sequences within species while driving differentiation between species (Arnheim, 1983). The coding regions are highly conserved, even between distantly related species, while non-coding DNA usually rapidly drifts apart, even between closely related species. Thus, using primers located in the conserved rDNA regions, variable regions can be amplified from a wide range of species in absence of prior sequence information. As such, the rDNA cluster has become an increasingly popular tool in molecular entomology (Collins & Paskewitz, 1996) and, in particular, as a means to develop diagnostic tests to differentiate cryptic Anopheline species. For example, diagnostic PCR assays based on segregating sequence variation in the ITS and/or IGS are now available to identify members of the *An. gambiae* Giles, *An. quadrimaculatus* Say, *An. punctulatus* Dönitz, *An. maculipennis* Meigen and *An. funestus* Giles complexes (Scott et al., 1993; Beebe & Saul, 1995; Cornet et al., 1996b; Proft et al., 1999;
Hackett et al., 2000). Furthermore, the D3 domain, known to be the most variable (coding) region in the 28S subunit provides a powerful tool for taxonomic studies in anopheles (Torres et al., 2000; Koekemoer et al., 1999).

Anopheles nili (Theobald) is widespread across tropical Africa (Hamon & Mouchet, 1961) and appears to be the major vector of malaria in some rural forested areas of central Africa, with entomological inoculation rates reaching 100 infectious bites/person/year (Carnevale et al., 1992). Larvae of An. nili are typically found in vegetation or in dense shade along the edges of streams and large rivers. The extensive morphological, ecological and ethological variations among An. nili populations have been reported by many authors (Gillies & De Meillon, 1968; Carnevale et al., 1992; Brunhes et al., 1999), suggesting that An. nili is a group of species.

To assess relevance of morphological characters as an accurate mean for classification within the group An. nili, we investigated sequence variation in the rDNA ITS2 and D3 domains of different morphological forms of this group. The analysis concerned the anthropophilic An. nili typical (T), the pale-winged An. nili Congo (Gillies & De Meillon, 1968) and three variants recently observed in Cameroon: An. nili A, An. nili B and An. nili Oveng. Our study included the recently described An. carnevalei (Brunhes et al., 1999) as well as the rare zoophilic and highly exophilic An. somalicus (Rivola & Holstein, 1957), characterized by slight morphological differences at the larval and pupal stages (Gillies & De Meillon, 1968; Gillies & Coetzee, 1987) while their adults resemble typical An. nili.

Experimental procedures

Mosquito collection and morphological identification

Specimens of An. nili s.l. were collected during surveys in four sites, Afan-Essokie (2°20’N 10°00’E), Mbebe (4°00’N, 11°02’E), Oveng (2°24’N, 10°22’E) and Simbock (3°55’N, 11°30’E), situated in the forested area of southern Cameroon. Specimens from Tibati (6°28’N, 12°37’E) and from Burkina Faso, Ivory Coast and Senegal were subsequently used to test for reliability of the PCR assay. Field sampling included both adult and larval collections.

Adult females were collected by landing catches on humans outdoors and indoors from 19.00 to 06.00 hours, and by pyrethrum spraying in selected bedrooms from 16.00 to 17.00 hours, during March and November. Adult specimens were identified morphologically in the field to separate An. nili sensu lato from other anopheline species, using morphological keys of Gillies & De Meillon (1968), Gillies & Coetzee (1987) and Brunhes et al. (1999). Anopheles nili s.l. specimens were then classified into six morphological types: An. nili typical, An. nili A, An. nili B, An. nili Congo, An. nili Oveng and An. carnevalei, according to the size and distribution of the pale fringe spots and pale veins spots of the wing. The typical (T) An. nili shows two pale spots on thealar fringe (CuA and M2 nerves). Variants A and B present one additional pale spot on thealar fringe at position (R4 + 5) nerve and CuA nerve, respectively, whereas the Congo form is characterized by the presence of both (R4 + 5) and CuA additional pale spots, and pale scales on the stem of the CuA nerve. A new morphological variant was sampled from the locality of Oveng (in South-western Cameroon) and will be hereafter referred to as An. nili Oveng. It is distinct from the Congo form by the presence of a large white base spot affecting the costa, with the CuA largely white. Finally, An. carnevalei differs from An. nili Congo by the presence of a large white pre-basal spot affecting slightly the costa and all the basal part of R1 nerve; the CuP is largely white and white spots are present on R3 and M1 veins (Brunhes et al., 1999).

Anopheles somalicus specimens were morphologically identified at the larval stage, as only aquatic stages of this species bears diagnostic morphological characters (i.e. longer clypeal hairs, see Gillies & de Meillon, 1968). Larvae of An. somalicus were found only in Mbebe.

DNA extraction, amplification and sequencing

Genomic DNA was extracted from whole larvae preserved in alcohol, single adult mosquitoes or from adult legs alone, following a slightly modified version of the protocol of Cornel & Collins (1996), and re-suspended in sterile water. ITS2 and D3 regions of the rDNA were amplified from approximately 10ng of template DNA in 25μl reaction mixture containing 2.5μL 10× reaction buffer (Qiagen, France), 1.5 mM MgCl2, 200 μM each deoxynucleotide triphosphate (Eurorgenec, Belgium), 0.625 U Taq DNA polymerase (Qiagen, France), 20 pmol each of ITS2a and ITS2b primers for ITS2 region, D3a and D3b primers for D3 domain.

ITS2a and ITS2b were designed from conserved position proximal to the 3′ and 5′ ends of the 5.8S and the 28S-rDNA, respectively (Collins & Paskewitz, 1996; Beebe et al., 1999). To amplify the variable D3 domain, D3a and D3b primers were designed in the 28S gene (Koekemoer et al., 1999; Sharpe et al., 1999). Sequences are as follows: ITS2a 5’TGTGAACCTGAGGACACAT3′ (forward) ITS2b 5’TATGCCTAATTTTCAGGGGTG3′ (reverse) D3a 5’GACCGTGCTTGGACACGG3′ (forward) D3b 5’TGGGAAGGACAGCTAAT3′ (reverse) PCR conditions included an initial denaturation step at 94°C for 3 min, followed by 35 cycles of 30s at 94°C, 30s at 55°C and 1min at 72°C with a final extension step of 10min at 72°C. After amplification, 5μL of the PCR products were analysed by electrophoresis onto a 1.5% agarose gel containing 0.5μg/mL ethidium bromide. Bands were revealed and photographed under UV light. After electrophoresis, the remaining PCR products were cleaned using spin columns and used for sequencing in both directions with the previous forward and reverse primers. Using CLUSTER V (Higgins & Sharp, 1988) we compared the sequences of complementary strands for each sample. Sequences were edited using the GCG package (Genetics Computer Group, 1997). Multiple alignment was performed with the PILEUP program of GCG. Genetic distances between haplotypes (Nei, 1987) were computed using MEGA 2.1 (available at http://www.megasoftware.net), under the pairwise deletion
option and using the Kimura 2-parameters correction to take into account multiple hits.

Original sequences of ITS2 and D3 have been deposited in the EMBL database under the following accession numbers: AJ429048 AJ429053 for *An. nili* typical, AJ429049 AJ429052 for *An. nili* Oveng, AJ438689 AJ438690 for *An. somalicus* and AJ429050 AJ429051 for *An. carnevalei*.

**Results**

**Analysis of D3 sequences**

The D3 domain of the 28S rDNA was successfully amplified in all specimens belonging to the seven morphological forms of *An. nili* using D3a and D3b primers. A single band of approximately 400 bp was revealed after electrophoresis. PCR products were directly sequenced in both directions using the same primers. The length of the sequenced fragments was 392 bp in *An. nili* typical, *An. nili* A, *An. nili* B and *An. nili* Congo and *An. carnevalei*, 390 bp in *An. somalicus*, and 397 bp in *An. nili* Oveng. At least eight sequences for *An. nili* typical, four sequences for each of *An. nili* A, *An. nili* B and *An. nili* Congo, eight sequences for *An. carnevalei*, five sequences for *An. somalicus* and six sequences for *An. nili* Oveng were aligned together. Four distinct haplotypes were observed among specimens morphologically identified as *An. nili* T (n = 8), three of which appeared in single copy in the dataset (haplotype diversity, h = 0.643; nucleotide diversity, π = 0.00098). The most common haplotype (five out of eight sequences, thus representing the consensus sequence for *An. nili* typical) was the only one observed in specimens belonging to *An. nili* A, *An. nili* B and *An. nili* Congo. Similarly, one single (specific) haplotype was observed among specimens of *An. nili* Oveng, *An. carnevalei* and *An. somalicus*, respectively. Sequence data obtained from the D3 domain are given in Fig. 1 and Table 1. Genetic distances between groups were thus computed through pairwise comparisons of consensus sequences (i.e. using one sequence per group). Pairwise genetic distances between haplotypes were low, ranging from 0.005 between *An. carnevalei* and *An. somalicus* to 0.024 between *An. nili* typical and *An. carnevalei* (Table 2).

**Analysis of ITS2 sequences**

PCR primers used to amplify the ITS2 region anneal to the 5.8S and the 28S coding regions of all species examined so far within the genus *Anopheles* (Porter & Collins, 1991). The ITS2 amplified products varied in size approximately between 500 bp and 600 bp within the *An. nili* group (data not shown). The banding pattern of ITS2 PCR products on agarose gels revealed four molecular entities but size differences were not sufficient to discriminate between forms. Sixty-two female individuals were sequenced: 30 *An. nili* typical, 5 *An. nili* A, 7 *An. nili* B, 3 *An. nili* Congo, 9 *An. carnevalei*, 5 *An. somalicus* and 3 *An. nili* Oveng. The length of the sequenced fragments was 450 bp for *An. nili* T, A, B and Congo, 480 for *An. carnevalei*, 513 bp for *An. somalicus* and 503 for *An. nili* Oveng. Figure 2 shows an alignment of the consensus sequences obtained for each of the seven morphological forms collected in Cameroon. As observed for the D3 domain, sequence comparison between forms revealed four distinct patterns of sequence variation, although low levels of variation were observed within each group (mean nucleotide diversity within group: π = 0.0057 ± 0.0009). Variation between haplotypes included nucleotide insertion-deletions and base substitutions. As shown in Fig. 2, *An. nili* typical and *An. nili* A, B and Congo presented highly homogenous consensus sequences. *Anopheles nili* Oveng had a microsatellite (AT) insertion at the 3’ end of the ITS2 fragment that was absent in the other forms. Pairwise genetic distances computed between consensus haplotypes are given in Table 2. Observed values were 7–20 times higher for this region than for the D3 domain and fixed differences between haplotypes provided the basis on which to develop specific PCR primers for an allele-specific PCR assay.

**Allele-specific PCR assay**

In order to achieve a quick, inexpensive and robust diagnostic tool for rapid identification of field-collected specimens down to their respective molecular form within the *An. nili* group, an allele-specific PCR assay (ASPCR) that combined five primers was developed. Fixed differences between haplotypes encountered in the ITS2 region were used as template to design diagnostic PCR primers (Fig. 2). The strategy we used for ITS2 allele-specific amplification is therefore similar to the approach of Walton et al. (1999) to distinguish between members of the *An. dirus* complex, and the approaches of Paskewitz & Collins (1990) and Scott et al. (1993) for sibling species identification within the *An. gambiae* complex.

A universal forward primer (ANU) that anneals to the 5’ end of the ITS2 of all *An. nili* s.l. specimens was designed,

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**Fig. 1.** D3 sequence from *An. nili* typical form.

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together with four reverse primers ANO, ANC, ANS and
ANT, specific to the An. nili Oveng (ANO), An. carnevalei
(ANC), An. somalisus (ANS) and to the typical (T) An. nili,
respectively, including variants A, B and Congo (ANT).
Primer sequences were defined so that at least four nucleo-
tides in their 3’ end would prevent hybridization to the
alternative haplotypes and guarantee specificity of the
assay. Sequences were as follows:
Primer ANU: 5’GATGCAACACATCTTCTGAGTG
CC3’
Primer ANO: 5’AGCAGCGTCACCTACGGTTCT
CC3’
Primer ANC: 5’CTGGTGGGTTCTTCTCTCT
CG3’
Primer ANT: 5’TGGCTGCTTTCTCGTGGGC
CCG3’
Primer ANS: 5’ATGCACAGGGGGTTC
CC3’

The Tm was 52°C, 56°C, 55°C, 56°C and 56°C, respect-
ively. Resulting PCR products differ from one another by at
least 30 bp, so that they are easily separated on regular agarose
gels. The size of the diagnostic band is 188 bp for the
typical (T) An. nili, 357 bp for An. nili Oveng, 408 bp for
An. carnevalei, and 329 bp for An. somalisus (Fig. 3). Hybrid
specimens were mimicked by mixing DNA from all combina-
tions prior to amplification. Two bands were obtained in
all cases at the expected size (Fig. 4).

Various PCR conditions were tested in order to optimize
amplification. PCR mixture consisted of 1.5 mM MgCl₂,
200 μM each dNTP (Eurogentec, Belgium), 2.5 μL 10×
Taq buffer, 0.625 U Taq polymerase (Qiagen, France) and
10 ng of template DNA in 25 μL final reaction volume. The
amount of each primer used in the PCR assay was 40 pmol
for ANU, and 10 pmol each for ANT, ANO, ANC and
ANS. PCR conditions included an initial denaturation
step at 94°C for 5 min, followed by 30 cycles of 30 s at
94°C, 30 s at 63°C and 1 min at 72°C with a final extension
step of 10 min at 72°C. The amplified fragments were separ-
ated by electrophoresis on a 2% agarose gel.

Validation of the assay on field-collected specimens

To test for the reliability of the ASPCR assay, a total of
226 wild An. nili s.l. adult females collected in five localities in
South Cameroon were investigated. PCR reactions using the
five primers ran on individual specimens produced the
expected pattern of size variation. The results confirmed
that all the 176 specimens morphologically identified as An.
nili typical, An. nili A, An. nili B, or An. nili Congo cor-
responded to typical (T) molecular form. The 30 specimens
identified as An. carnevalei based on morphological char-
ters unambiguously showed an An. carnevalei molecular pat-
tern and all of the 20 specimens identified as An. nili Oveng
belonged to the Oveng molecular form. Field-collected An.
somalisus larvae (n = 15) and emerging adults (n = 10) of An.
somalisus were correctly identified. It is noteworthy that none
of the adult specimens An. nili s.l. tested showed the An.
somalisus-specific PCR product. Moreover, five specimens
Table 2. Pairwise genetic distance between consensus haplotypes of the rDNA ITS2 and D3 regions from south Cameroon populations of the *An. nili* group. Alignment gaps were excluded from the analysis under the pairwise-deletion option and the Kimura 2-parameters correction was applied to take into account multiple hits. Figures above the dashed line are ITS2, figures below the dashed line are D3.

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<td><em>An. nili</em> B</td>
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identified as *An. nili* collected in Burkina Faso, Ivory Coast and Senegal, were tested. All of them belonged to *An. nili* typical (T) molecular form. Sensitivity of the assay therefore was 100% and no hybrid pattern between members of the *An. nili* group was observed.

In tropical Africa, members of the *An. nili* group are usually sympatric with *An. funestus*, members of the *An. gambiae* complex and *An. moucheti* Evans among others. In the conditions of our technique, no PCR product was obtained with 10 specimens of each of these other sympatric species.

Discussion

Distinction between members of the *An. nili* group is presently based mainly on morphological characters. However, these criteria are often difficult to apply because of a number of biological and/or technical issues (e.g. shared overlapping characters, inadequate sampling and preservation of specimens), as for example in Cameroon where seven different morphological forms were observed. Hence, the taxonomic status of each of the reported morphological variants of *An. nili* s.l. remained largely subjective. Precise knowledge of the biology, and thus of the specific importance of members of the group as malaria vectors, has been limited by the absence of reliable diagnostic characters. Our study of sequence polymorphism of the ITS2 and D3 domains of rDNA has overcome this limitation by developing an allele-specific PCR assay which can serve as a reproducible and standardized diagnostic tool for distinguishing between members of the *An. nili* group. Similar approaches have provided powerful diagnostic tools for the study of other anopheline species complexes, such as *An. maculipennis* (Porter & Collins, 1991), *An. gambiae* (Paskewitz et al., 1993), *An. quadrimaculatus* (Cornel et al., 1996) and *An. funestus* (Kockemoer et al., 2002).

Sequence variation in both the rDNA ITS2 and D3 domains among the seven morphological forms of *An. nili* s.l. collected in Cameroon defined four genetic clusters. The first cluster included specimens morphologically identified as typical *An. nili*, variants A and B and the Congo form, whereas the three others included specimens identified as *An. carnealet*, *An. nili* Oveng and *An. somalicus*, which consistently showed unique ("specific") haplotypes. These results are in partial agreement with morphological features, and suggest that the *An. nili* group may comprise at least four reproductively distinct biological species. Indeed, genetic distances between ITS2 consensus haplotypes of each of these molecular forms were in the range 0.11–0.25, values higher than expected between conspecific populations (Avise, 1994), and similar to those observed within the *An. funestus* group (Hackett et al., 2000; Kockemoer et al., 2002), between members of the North American *An. quadrimaculatus* complex (Cornel et al., 1996) and between the isomorphic South American species *An. trinkai* and *An. dunhami* (Lounibos et al., 1998). To date, however, specimens of the different molecular forms of the *An. nili* group have only been found in allopatry, thus preventing further interpretation of their taxonomic status. Putative areas of sympathy are actually being screened and multilocus genetic analyses are under way, using isozymes as well as recently developed microsatellite markers, to ascertain the hypothesis of speciation within the *An. nili* group.

As mentioned above, owing to the limitations of morphological identification methods, very few data are available on biology of the larvae and adults, on the distribution and on the exact role in malaria transmission of the different members of the *An. nili* group. The multiplex PCR that we have developed is simple, sensitive, inexpensive and usable in a laboratory having basic molecular equipment. It can be used on all stages of development, on both sexes (although hybrid males may be impossible to identify if all rDNA repeats were located on the X chromosome, as is the case for *An. gambiae*, see Collins et al., 1989) and correct amplification has been obtained from dry as well as alcohol preserved specimens. Moreover, as frequently highlighted as an advantage of PCR-based methods, only a small quantity of biological material (for example a single leg) would contain sufficient template DNA to allow amplification, which leaves the possibility of using the remaining body of the mosquito, even alive, for other studies. Additionally, the technique allows simultaneous testing of a great number of specimens, as usually required during epidemiological investigations. *Anopheles nili* s.l. mosquitoes from Senegal, Burkina Faso, Côte d’Ivoire and Cameroon were successfully identified using this protocol, suggesting that the technique
Fig. 2. Alignment of ITS2 consensus sequence from isolates representing each member of the An. nili group. The sequences in the alignment represent An. nili typical form (TFO), Congo form (CFO), A form (AFO), B form (BFO), An. somalicus (SOM), An. carnevalei (CAR), and Oveng form (OVE). The dots (·) indicate that the sequence at that point is the same as in An. nili Congo. Dash (–) indicates alignment space. Areas from which the primer where designed are underlined. The primers are as follows: ANU (universal primer), ANT (An. nili typical, Congo, A and B), ANS (An. somalicus), ANO (An. nili Oveng) and ANC (An. carnevalei).
Fig. 3. Photograph of 2% agarose gel showing the amplification products from a single PCR containing the universal primer and the four specific primers. Lane M is 100-bp DNA size marker, lanes 1 = *An. nili* typical, 2 = *An. nili* congo, 3 = *An. nili* A, 4 = *An. nili* B (188 bp), 5–6 = *An. somalicus* (329 bp), 7–8 = *An. nili* Oveng (357 bp) and 9–10 = *An. carnevales* (408 bp). Lane T is a negative control.

Fig. 4. Ethidium bromide-stained 2% agarose gel showing the PCR products from different molecular form in *An. nili* group and mixtures of DNA to simulate hybrids. Lane M: 100 bp DNA size marker, 1 = *An. carnevales*; 2 = *An. nili* Oveng; 3 = *An. somalicus*; 4 = *An. nili* typical; 5 = *An. carnevales* plus *An. nili* typical; 6 = *An. carnevales* plus *An. somalicus*; 7 = *An. carnevales* plus *An. nili* Oveng; 8 = *An. nili* Oveng plus *An. nili* typical; 9 = *An. somalicus* plus *An. nili* typical; 10 = *An. nili* Oveng plus *An. somalicus*. Lane T is a negative control. All reaction conditions and the size of PCR products were as described in the text.

has a continent-wide application. Routine application of this new diagnostic test to complement morphological identification (not to be neglected), should help to increase our knowledge of malaria transmission dynamics in areas with vector populations of the *An. nili* group, and contribute to better targeting and implementation of specific and selective strategies for malaria vector control in Africa (Hougaard et al., 2002).

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