# A polymerase chain reaction assay to determine infection of Aedes polynesiensis by Wuchereria bancrofti

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The sensitivity of a previously described polymerase chain reaction (PCR) assay was improved to detect a single mosquito, infected by as few as 1-2 microfilariae of Wuchereria bancrofii, among 20-50 uninfected mosquitoes. Wild-caught Aedes polynesiensis were used to compare assessment of infection by dissection of individuals with the PCR assay of pools of mosquitoes. The PCR assay was at least as sensitivite as dissection for detection of mosquitoes infected with W. bancrofti.

Keywords: filariasis, Wuchereria bancrofti, Aedes polynesiensis, polymerase chain reaction

#### Introdction

At least 120 million people are thought to be infected by Wuchereria bancrofti and, to a lesser extent, by Brugia malayi or B. timori (see Ottesen & RAMACHANDRAN, 1995). Evaluation of chemotherapy campaigns is based not only on detection of the parasite or specific parasite antigens in patients but also on monitoring the infection of mosquito populations. Traditionally, this has involved the dissection and microscopical examination of hundreds to thousands of mosquitoes, which is very time consuming. Another drawback of the dissection method is the risk of confusing the several filarial spe-

cies which may co-infect a single mosquito.

The development of deoxyribonucleic acid (DNA) probes and polymerase chain reaction (PCR) assays for detection of parasites has been recently reviewed (WEISS, 1995). Development of molecular techniques to detect filarial parasites in pools of vectors is an objective of the World Health Organization (WHO, 1993). A preliminary PCR assay based on the amplification of a DNA repeat sequence from W. bancrofti, the 'SspI DNA repeat', and using specific primers has been described by CHAN-TEAU et al. (1994). In this assay, a 188 base pair (bp) DNA fragment was amplified specifically from W. bancrofti but not from DNA of other filariae, humans or mosquitoes; this fragment is present in all the geo-graphical isolates of W. bancrofti so far tested. By contrast, it is not stage-specific.

A PCR method reported by CHANTEAU et al (1994) detected a single third-stage larva (L3) of W. bancrofti added to a pool of 50 heads of Aedes polynesiensis, the main vector of bancroftian filariasis in French Polynesia. However, for convenience, the PCR needs to be performed on pools of whole mosquitoes instead of heads. A PCR assay must also be able to detect any infected mosquito, even by a single parasite, in pools of mosquitoes. With these conditions fulfilled, it should be possible to examine large numbers of mosquitoes in a much shorter time than by dissection. Although the DNA probe considered here is not stage-specific, evaluation of infection rates in mosquito populations, instead of infectivity rates, would be of great help in monitoring chemother-

apy campaigns.

We have improved the PCR assay described by CHAN-TEAU et al. (1994) so that it will now detect a single L3 in a pool of 50 to 100 whole A. polynesiensis or a single mosquito, infected by about 2 microfilariae (mf) in 20 to 50 whole mosquitoes. Furthermore, we have shown that the PCR assay performed on pools is at least as sensitive as dissection in determining infection rates of wild popula-

tions of A. polynesiensis.

#### Materials and Methods

Polymerase chain reaction assay

The extraction of DNA from heads or whole mosqui-

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toes was modified from the method of CHANTEAU et al. (1994) as follows. Mosquitoes were dried for 3 h at 90°C, then separated heads or whole insects were carefully crushed with a sterile pestle tissue grinder in a 1.5 mL plastic tube. The crushed material was washed twice with 1 mL of 0·1 M NaCl, 30 mM Tris-HCl (pH 7·8), 30 mM ethylenediaminetetraacetic acid (EDTA), 10 mM 2-β-mercaptoethanol, and 0.5% Nonidet P40®. The supernatant was discarded and DNA was released by lysing the pellet with 100 μL of 0·1 M NaOH and 0·2% sodium dodecyl sulphate for 1 h at 37°C (or 200  $\mu$ L if the number of mosquitoes was above 10), neutralized with 5 (or 10)  $\mu$ L of 2N Hcl, and then mixed thoroughly with 1 mL of 4.5M guanidine hydrothiocyanate, 50 mM Tris-HCl (pH 6.4), 1.2% Triton X100®, and 20 mM EDTA. DNA was precipitated by addition of 40  $\mu$ L of silica particle suspension (Sigma) and incubation for 10 min at room temperature. Following centrifugation for 10 s at 12 000 g in a Microfuge®, the silica particles were washed twice with 4.5M guanidine hydrothiocyanate and 50 mM Tris-HCl (pH 6.4). The silica pellet was then dried at 56°C for 10 min with the tube cap removed, and the beads were resuspended in 100  $\mu L$  of TE buffer (10 mM Tris-HCl [pH8], 1 mM EDTA) and incubated at 56°C for 10 min to elute the DNA. After 10 s centrifuga-

tion, the supernatant was kept for PCR. The PCR reaction was performed on 1 to 10  $\mu$ L of mosquito supernatant in a final volume of 50  $\mu$ L, and included 2 units of *Taq* polymerase (Promega), 400 pM of both primers, NV1 and NV2 (CHANTEAU et al., 1994), and  $2\bar{0}0 \,\mu\text{M}$  of each dideoxynucleotide in 50 mM KCl,  $1\bar{0}$ mm Tris-HCl (pH9), 0.1% Triton X100®, and 1.5 mm MgCl<sub>2</sub>. The temperature programme for the PCR was 5 min at 92°C, then 30 cycles of 15 sec at 92°C, 1 min at 55°C, 1 min at 72°C, and a final 10 min at 72°C.

Ten  $\mu$ L of the PCR product were loaded on to a 1.5% agarose gel and a unique band of 199 bp was visualized by ethidium bromide staining. A negative control for the PCR assay, using water instead of DNA extract in the reaction mix, was included with all runs. A positive control was also included, consisting of 1 pg (or 0·1 pg) of W. bancrofti mf genomic DNA, which corresponds to c. 1% of an L3 larva per  $\mu$ L.

Because A. polynesiensis is also the vector of Dirofilaria immitis, the dog heartworm, in French Polynesia, we checked the specificity of the PCR assay for W. bancrofti. The sensitivity limit for the assay was 0.1 pg of W. bancrofti DNA, but no comparable DNA fragment was amplified from up to 800 pg of DNA from a local strain of  $oldsymbol{D}$ . immitis.

Sensitivity and reproducibility of the assay to detect W. bancrofti *L3* 

W. bancrofti L3 were obtained from laboratory-bred A. polynesiensis fed on a microfilaraemic Polynesian volunteer. Fourteen days after blood-feeding, the mosquito heads were cut off and immersed in RPMI-1640 medium to allow passive release of L3, which were then further purified by filtration through fine mesh gauze. DNA extraction and the PCR were carried out on pools of 50 or 100 heads of uninfected A. polynesiensis or on whole mosquitoes 'spiked' with one L3.

Our ability to detect L3 within whole mosquitoes by the PCR assay was tested with a batch of A. polynesiensis, 90% of which were infected with a mean of 6 L3 (range 1–10). Ten pools of 50 mosquitoes containing either 1, 2 or 3 putatively infected individuals were examined.

Negative controls for the extraction procedure were included in all experiments to avoid false positive PCR results. Every tenth extraction of mosquito heads or whole mosquitoes was done using uninfected laboratory-bred mosquitoes. The extracts from these negative controls were then included in the PCR assay.

Sensitivity of the assay for detecting microfilariae in laboratory-bred mosquitoes

The potential of the PCR assay to detect A. polynesiensis infected with mf was determined as follows. A batch of A. polynesiensis infected with a mean of about 2 mf each was obtained by artificial blood feeding for 1 h on 3 mL aliquots of human blood containing c. 1000 mf/mL (FAILLOUX et al., 1991). After the blood meal, fed mosquitoes were separated and frozen at -20°C.

The number of mosquitoes containing mf, as determined by dissection of 30 individuals, was compared with the detection rate by PCR using pools of mosquitoes. The sensitivity of PCR was also investigated using 30 pools of 5, 20 and 50 uninfected mosquitoes 'spiked' with a single putatively mf-infected mosquito to determine the effect of pool size on the sensitivity of the as-

Comparison of PCR and dissection of wild-caught mosquitoes A. polynesiensis were collected in 2 distinct geographical areas of Tahaa Island (Society Archipelago, French Polynesia), where the human populations had been previously treated with either ivermectin at 400 μg/kg (IVR area) or a combination of ivermeetin at 400 µg/kg plus diethylcarbamazine at 6 mg/kg (IVR+DEC area) (MOULIA-PELAT et al., 1995). Approximately 2000 daybiting A. polynesiensis were collected 2, 4 and 6 months after the treatment in the IVR area and 2 and 6 months after in the IVR+DEC area, by uninfected persons acting as both bait and collectors. The mosquitoes were then divided into 2 groups for each drug and time combination. Mosquitoes (>1300) from one group were individually dissected for identification of filaria larvae and developmental stages as described by LARDEUX et al. (1995). The other group was killed by freezing within 2-4 h after capture and stored at -20°C until used in PCR assays. In this field study, we examined 50 pools of 5 mosquitoes from each mosquito group to determine the proportion of pools which was positive by PCR.

The PCR results from pools were compared with the results of dissection of individual mosquitoes as follows. Using the real numbers of A. polynesiensis dissected and infected as input (e.g., 1369 and 6, Table 2), we randomly sampled by computer 50 groups of 5 among the dissected mosquitoes. A group containing at least one infected mosquito was considered as positive. This allowed us to determine the number of positive groups among 50 in one sampling. This simulation was repeated 1000 times to determine the number  $N_i$  and the probability  $P_i$  (=  $N_i/1000$ ) of finding 0, 1, 2, ... i positive pools. The simulated number of positive pools detected by dissection ('dissection positive pools') was the product of  $P_i \times i$  and was compared with the number of pools positive by PCR ('PCR positive pools').

Statistical analysis

The  $\chi^2$  test was used to test the significance of differences (P<0.05) in proportions of positive individuals or pools in the detection of mosquitoes infected with mf and in the assay with wild-caught mosquitoes. In this

latter experiment, a t test for paired observations (P<0.05) was also used to compare the results of the 2 methods.

#### Results

Sensitivity and reproducibility of the PCR assay to detect W. bancrofti L3

All 10 pools of either 50 or 100 heads 'spiked' with one L3 were positive by PCR. Using whole mosquitoes instead of heads, all 10 extracts from pools of 50 mosquitoes were also positive by PCR, while only 5 of 10 pools of 100 mosquitoes were positive.

The PCR assay detected 8 of 10 pools of 50 mosquitoes containing a single putatively infected mosquito (from a batch of mosquitoes 90% infected with a mean of 6 L3 per infected mosquito). When the number of putatively

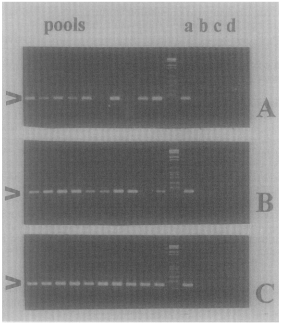


Figure. Detection of *W. bancrofti* in *A. polynesiensis* by PCR assay of 10 pools of 50 whole mosquitoes. A: 49 uninfected mosquitoes + 1 putatively infected; B: 48 uninfected mosquitoes + 2 putatively infected; C: 47 uninfected mosquitoes + 3 putatively infected. Controls used as templates in PCR assays: a, 1 pg *W. bancrofti* DNA; b, 0·1 pg *W. bancrofti* DNA; c, water; d, DNA from 50 uninfected *A. polynesiensis*. Arrow heads indicate the 188 base pair product.

Table 1. Comparison of dissection and PCR for the detection of A. polynesiensis experimentally infected with W. bancrofti microfilariae

No. of mosquitoes per sample	No. of infected mosquitoes/pool
Dissection	
30 individual blood-feda	
Examination no. 1	24/30 <sup>b</sup>
Examination no. 2	25/30
PCR	
30 individual blood-fed	27/30
30 pools of 1 blood-fed+4 uninfected	28/30
30 pools of 1 blood-fed+19 uninfected	25/30
30 pools of 1 blood-fed+49 uninfected	l 16/30 <sup>b</sup>

<sup>&</sup>lt;sup>a</sup>The 30 individually dissected mosquitoes were examined twice and contained a mean of 1.9 microfilariae per infected mosquito (SE=0.32, range 1-7) at the first examination and a mean of 2.4 microfilariae per infected mosquito at the second examination (SE=0.37, range 1-8).

bSignificantly different ( $\chi^2$ , P < 0.05).

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Table 2. Comparison of dissection and PCR as a method to estimate W. bancrofti infection rates in wild-caught A. polynesiensis

	Drug treatment <sup>a</sup>					
	IVR	IVR	IVR	IVR+DEC_	IVR+DEC	
Months after treatment	2	4	6	2	6	
No. of mosquitoes						
Dissected	1369	1815	1774	1635	1944	
Infected	6 (0.44%)	15 (0.83%)	24 (1.35%)	11 (0.67%)	6 (0.31%)	
Proportion of positive mos	squito pools			, ,	` ,	
Dissectionb	1.1/50	1.9/50	3·1/50	1.6/50	0.7/50	
PCR	5/50	7/50	7/50	4/50	6/50	
$\chi^2$ (d.f.=1)	1.47	2.07	0.93	0.37	2.78	
P	0.22	0.15	0.34	0.54	0.09	

<sup>&</sup>lt;sup>a</sup>Drug used to treat population in the area where mosquitoes were caught 2, 4 or 6 months later; IVR=ivermectin (400 μg/kg), DEC=diethylcarbamazine (6 mg/kg).

infected mosquitoes was increased to 2 or 3, the PCR detected respectively 9 and 10 of 10 pools (Fig. 1).

Sensitivity of the assay to detect mf in laboratory mosquitoes

Dissection and 2 microscopical examinations of 30 mosquitoes artifically fec on blood containing W. bancrofti mf indicated that 78% of these mosquitoes were infected by an average number of 2·1 mf/mosquito. As shown in Table 1, infection rates determined by dissection and PCR of individual mosquitoes were not significantly different ( $\chi^2 = 1.17$ , df=1, P=0.23).

Moreover, the sensitivity of the PCR assay to detect a single infected mosquito was not affected if this mosquito was mixed with pools of up to 20 uninfected mosquitoes. A significant loss of sensitivity was observed in pools of 50 mosquitoes (Table 1,  $\chi^2=4.80$ , df=1, P<0.02).

Comparison of PCR and dissection of wild-caught mosquitoes

Infection rates of wild-caught mosquitoes ranged from 0.3 to 1.35%. The numbers of 'dissection positive pools', calculated by simulation analysis from the results of individual dissections, were always lower than the numbers of pools which were positive by the PCR assay (PCR positive pools') (Table 2). The difference between dissection positive pools' and PCR positive pools' was significant  $(t_4=7.92, P<0.001)$ . However, the differences between proportions of 'dissection positive pools and 'PCR positive pools', for each time and drug combination, were not significantly different ( $\chi^2$  test) (Table 2).

# Discussion

In this study we improved a PCR assay based on the amplification of a DNA repeat sequence from W. ban-crofti, the 'SspI DNA repeat'. We can now detect a single infected whole mosquiot among a pool of as many as 20 to 50 uninfected mosquitoes.

Applied to wild-caught mosquitoes, the assay was shown to be at least as sensitive as dissection. We found higher rates of infection by PCR than by dissection in the 5 groups of mosquitoes examined. Two points must be considered. First, the dissection of several hundreds of mosquitoes was carried out over a short period of time by several workers on populations of mosquitoes infected with few larvae. Infected mosquitoes harboured a mean number of 3.9 L1, 2.3 L2, 3.1 L3 (F. Lardeux, unpublished observations). Detection of L3 is relatively easy. However, detection of the smaller stages (L1) is more difficult as it requires accurate dissection of mosquito thorax muscles to release the parasites and render them visible. It is therefore likely that some mosquitoes infected by these smaller stages might have been missed. By contrast, extraction of DNA from mosquitoes is very efficient because we have shown that a mosquito in-

fected by only 1-2 mf is detectable by PCR. In the field, infected host-seeking females will not usually contain mf, only the later stage larvae which, being larger, are even easier to detect than mf. Therefore, the method should be more sensitive in practice than it proved to be in this experimental study. Secondly, we should also investigate the possible persistence of DNA from W. bancrofti parasites which do not complete development within the mosquito but which might still be amplified by PCR, leading to false positive results.

We are currently working to improve the PCR assay to measure accurately the amount of DNA amplified from infected mosquitoes. Although the DNA fragment amplified in the SspI PCR assay is not stage specific, it has the great advantage of being specific for W. bancrofti and ubiquitous in all the W. bancrofti isolates so far tested. At present, this assay estimates the number of pools of mosquitoes which contain at least one infected mosquito. This is of practical interest in areas where infection rates are very low. With the present state of technology, a technician working 8 h per day with a single thermocycler can test about 30-40 pools of mosquitoes (i.e. 1500 to 2000) per day. This is to be compared with the ability of a well trained technician to dissect and examine about 100-200 mosquitoes a day. The PCR assay also allows the screening of mosquito extracts for the presence of other infectious agents, using different molecular probes.

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# Note added in proof

A computer program (Poolscreen<sup>TM</sup>) which predicts infection rates of populations based upon screening pools was published while this paper was in press (KATHOLI, C. R., TOÉ, L., MERRIWHEATHER, A. & UNNASCH, T. R. (1995). Determining the prevalence of *Onchocerca volvulus* infection in vector populations by polymerase chain reaction screening of pools of black flies. *Journal of Infectious Diseases*, 172, 1414–1417). Using the Poolscreen<sup>TM</sup> program, the infection rates (and 95% confidence limits) of mosquito populations calculated from our PCR on pools of mosquitoes (Table 2) were 2·09% (0·67–4·80), 2·97% (1·19–6·03), 2·97% (1·19–6·03), 1·65% (0·44–4·18) and 2·52% (0·92–5·41) respectively for groups IVR month 2, IVR month 4, IVR month 6, IVR+DEC month 2 and IVR+DEC month 6. In this paper we calculated the numbers of 'dissection-positive pools' by simulation analysis using data from individual dissections. The equivalent calculations using the Poolscreen<sup>TM</sup> program adequately fit our data.

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