Determination of the prevalence of lymphatic filariasis among migrant workers in Kuwait by detecting circulating filarial antigen

Jamshaid Iqbal1 and Ali Sher2

The main objective of this study was to determine the prevalence of filarial infection among migrant workers in Kuwait. The study was conducted from April 2000 to November 2003. A total of 1050 migrant workers (> 90% from the Indian subcontinent) from filarial endemic countries and 260 individuals residing in Kuwait as controls (50 healthy Kuwaiti blood donors, 50 microfilaria-negative individuals from endemic areas and 160 patients with other parasitic infections) were screened for filarial infection. All specimens were tested for microfilaraemia by microscopy of nucleopore-filtered blood (NFB) and detection of circulating filarial antigen (CFA) by an immunochromatographic test (ICT) and the TropBio assay. The overall prevalence of filarial antigenaemia was 18 ± 3% (192 individuals) using the ICT and 20 ± 3% (213 individuals) using the TropBio assay. Thirty-two cases (3%) of *Wuchereria bancrofti* were detected by microscopy and the mean microfilaria count in these cases was 816 microfilariae ml⁻¹. CFA was detected only in two of the 260 control subjects. Statistical analysis to calculate the sensitivity, specificity and prevalence of infection was carried out using maximum-likelihood statistical methods. The overall sensitivity and specificity of the ICT and TropBio assay to detect CFA were comparable. Compared with NFB microscopy, the sensitivity of the ICT was 93 ± 8% and specificity ranged from 84 to 100%. The sensitivity and specificity of the TropBio assay were 90 ± 1 and 100%, respectively. However, the ICT failed to detect CFA in two cases with a microfilarial load of < 20 microfilariae ml⁻¹. In conclusion, the prevalence of filarial infection among the migrant workers in Kuwait was 18 ± 3% as determined by the ICT.

**INTRODUCTION**

Lymphatic filariasis continues to be a major source of permanent disability and an impediment to socio-economic development in 73 countries where more than 1 billion people are at risk and over 120 million are infected. *Wuchereria bancrofti* is responsible for approximately 90% of infections (World Health Organization, 1999).

Early and efficient diagnosis of *W. bancrofti* infection is a key step in monitoring, treatment and subsequent eradication of lymphatic filariasis. Microscopic examination of a thick blood film or nucleopore-filtered blood (NFB) to detect microfilariae (mf) lacks sensitivity and provides only an indirect measure of the adult worm load. Moreover, the current filarial antibody detection tests remain unable to distinguish between active and past infections (Bruschi & Castagna, 2004).

Recently, an immunochromatographic test (ICT) using whole-blood samples was introduced. The test detects soluble circulating filarial antigen (CFA) in blood (McCarthy, 2000; Weerasooriya et al., 2003). The ICT has been evaluated for its sensitivity and specificity in various epidemiological settings and the reported sensitivity ranges from 73 to 100% (Braga et al., 2003; Chandrasena et al., 2002; Omar et al., 2000; Reddy et al., 2004).

No indigenous case of filarial infection in Kuwait has been reported as yet; however, the presence of *Culex quinquefasciatus*, the vector of bancroftian filariasis, and a large number of potentially infected individuals in Kuwait may initiate local transmission. Every year, more than 45 000 migrant workers arrive in Kuwait from areas where filarial infection is endemic. On arrival in Kuwait, these workers are screened for filarial infection, malaria, hepatitis, human immunodeficiency virus and tuberculosis infection before they are issued a resident permit. Screening for filarial infection is done by microscopy of a thick blood film. Due to technical, logistical and administrative constraints, these
workers can only be screened during the daytime, although mf are present mostly in the blood circulation at night. This study was carried out to determine the level of filarial infection among migrant workers in Kuwait by detecting filarial antigen using the ICT and the TropBio assay.

METHODS

**Study population.** The study was performed at the Ports and Borders Center, Ministry of Health, Kuwait, and the Department of Microbiology, Faculty of Medicine, Kuwait University, during the period April 2000 to November 2003. A total of 1050 immigrant workers from countries where filarial infection is endemic were selected randomly and enrolled in the study. The majority (>95%) of these individuals were from India (Andhra Pradesh, Bihar, Tamil Nadu and Uttar Pradesh – the filaria belt), the Philippines and Sri Lanka. The others were from Tanzania, Mali, Gambia and Sudan. For comparison, control blood specimens were also collected from 260 individuals residing in Kuwait comprising 50 healthy Kuwaiti blood donors, 50 individuals originally from endemic areas who had been living in Kuwait for more than 10 years and had tested negative repeatedly for mf, and 160 patients with other parasitic infections (hydatid disease, amoebiasis, malaria, schistosomiasis) from areas non-endemic for filaria. Socio-demographic information and their history of travel were collected from all individuals enrolled in the study.

All specimens were tested blindly at the Ports and Borders Center, Kuwait, and the Department of Microbiology, Faculty of Medicine, Kuwait University. All participants were informed about the objectives of the study and their consent was obtained. The Ethical Committee of the Faculty of Medicine approved the study.

**Specimen collection.** Due to administrative and logistical reasons, all blood specimens were collected between 8 a.m. and 10 a.m. Blood (2 ml) was collected into an EDTA-coated tube and tests were performed on the same day. Blood specimens were also collected at midnight from 25 microscopy-negative but ICT-positive individuals for retesting of filtered blood by microscopy.

**NFB microscopy.** The blood was diluted with 9 ml 2% formalin in normal saline and filtered using a nucleopore polycarbonate membrane (pore size 5 μm, diameter 25 mm). The filter membranes were stained with Giemsa stain and examined by two experienced microscopists under low-power magnification for the presence of mf.

**ICT.** The ICT uses paired antibodies (polyclonal and monoclonal antibodies) specific for *W. bancrofti* antigen (AD12.1) conjugated to visible colloidal gold particles and immobilized on the test strip. The test was performed as described by the manufacturers (NOW ICT Filariasis Test; Binax). Briefly, 50 μl whole blood was added to the sample pad, allowing any filarial antigen present in the plasma to bind to the colloidal gold-labelled antibody on the strip. Cards exhibiting the control line were considered valid. Reading and interpretation of the results followed the manufacturer’s instructions.

**TropBio ELISA assay.** The TropBio ELISA assay (ICI Tropical Biotechnology) detects and quantifies *W. bancrofti* antigen in blood specimens. The assay was performed as described previously (Weerasooriya et al., 2003). Briefly, each specimen was tested in duplicate. The mean of the results was classified against seven known standard concentrations of *Onchocerca gibsoni* antigen and a standard curve was produced relating absorbance and antigen content. The antigen concentration and the antigen titre group for each specimen were determined from the standard curve. A cut-off value of 32 antigen units (AU) (titre group 4) was considered positive for CFA and a value of >8000 AU was assigned to titre group 7.

**Data analysis.** The sensitivity of NFB microscopy and the ICT is unknown, as many individuals may harbour adult worms without having mf in their peripheral blood. Therefore, we assumed that both NFB microscopy and the ICT might be 100% specific, but not 100% sensitive. Making this assumption, the sensitivity and specificity of the NFB microscopy and the ICT and the prevalence of filarial infection among the immigrant workers were calculated using statistical maximum-likelihood (ML) methods (Goldberg & Wittes, 1978) and the specificity by using the equation derived by Staquet et al. (1981).

**RESULTS**

The presence of filarial infection was examined in 1050 migrant workers [603 (57.4%) males and 447 (42.6%) females aged between 14 and 61 years (mean ± SD 29.2 ± 3.2)] by NFB microscopy and for the presence of CFA using the ICT and the TropBio assay. The overall prevalence of filarial antigenaemia in this group was 18.3% (192 individuals) using the ICT and 20.3% (213 individuals) using the TropBio assay (Table 1). However, microfilaremia was present only in 34 cases (3%); 19 cases had <100 mf ml\(^{-1}\), 10 cases had 100–750 mf ml\(^{-1}\) and five cases had >750 mf ml\(^{-1}\). The mean mf count in these cases was 816 mf ml\(^{-1}\) (range: 17–1283 mf ml\(^{-1}\)). Only two cases of *Brugia malayi* were observed; both of these cases were from the Philippines. These two cases were excluded from further analysis by the ICT and the TropBio assay. All other cases had *W. bancrofti* infections and were also positive for CFA using the ICT and the TropBio assay. Given the logistical constraints of collecting midnight blood specimens from all subjects, a small group of 25 NFB-negative, ICT-positive individuals was retested by taking their blood specimens at night. Only two of these 25 individuals showed mf in NFB in their midnight specimens; the density of mf in these two cases was <50 mf ml\(^{-1}\) (data not shown). Only two of the 260 control specimens were positive for CFA using the ICT and the TropBio assay (Table 1). Both of these cases were males from Andhra Pradesh, India, working in Kuwait and had not travelled to India in the last 5 years. These cases were excluded in the determination of test specificity. The majority of cases (>90%) with antigenaemia and microfilaremia were males in the 24–41 year age group.

The performance of the ICT in detecting antigenaemia was compared with NFB microscopy (Table 2). Among 192 cases with CFA, 30 cases (15.6%) had microfilaremia. However, two mf-positive cases with a density of <20 mf ml\(^{-1}\) did not show any detectable CFA (Table 2). The sensitivities of NFB microscopy and the ICT and the prevalence of infection among the migrant workers were calculated using the ML statistical method. The prevalence of filarial infection in the migrant population was estimated to be 19.5%; the sensitivity of NFB microscopy was only 15.9%, whilst the sensitivity of the ICT was 93.8%. The specificity of the ICT ranged from 84.1 to 100%, based on the prevalence of infection using NFB microscopy and the ICT, calculated as described in Methods.

The TropBio assay gives a quantitative response expressed in AU or titre groups. Of 213 cases showing various levels of
CFA, 75 had CFA levels of 32–128 AU (group 4), 48 had CFA up to 512 AU (group 5), 71 had CFA up to 2048 AU (group 6) and 19 had CFA > 8000 AU (group 7). Analysis of the relationship between CFA levels and mf density showed a positive association, but was not statistically significant (P > 0.05).

The performance of the TropBio assay for detection of CFA showed a similar pattern to that of the ICT (Table 2). The ICT detected CFA in 192 (90.1%) of the 213 cases that were positive with the TropBio assay. Comparing these two tests, the sensitivity of the ICT was 92.1% (Table 2). Furthermore, the overall sensitivity of the ICT and the TropBio assay for detection of CFA in cases with different mf densities was comparable.

**DISCUSSION**

The recent development of reliable tests for detection of CFA in *W. bancrofti*-infected individuals has been highly successful. Several studies have reported the sensitivity, specificity and predictive values of the ICT in different epidemiological settings (Braga *et al.*, 2003; Chandrasena *et al.*, 2002; Omar *et al.*, 2000). In this study, the sensitivity of the ICT was 93.8%. Contrary to most previous studies, the performance of the ICT, the TropBio assay and NFB microscopy was determined in all study subjects in this study. The imperfect ‘gold standards’ like NFB microscopy generally introduce bias, such as an underestimation of the prevalence of filarial infection, specificity and the positive predictive value of the new test. Therefore, in this study, all statistical analysis was done using ML methods and Staquet’s equation (Goldberg & Wittes, 1978; Staquet *et al.*, 1981); these are considered to be the ideal methods for calculation of sensitivity and specificity of new screening assays.

The overall prevalence of filarial antigenaemia among migrant workers was 18.3% and microfilaraemia was present only in 3% of cases. Since mf are present mostly in the blood circulation at night, the number of microfilaraemic individuals detected using daytime blood samples was expected to be significantly lower than that obtained with night-time blood samples. However, as mentioned above, night-time blood samples were not available due to unavoidable administrative and technical reasons. Given the logistical

<table>
<thead>
<tr>
<th>Categories of study cases</th>
<th>No. tested (males/females)</th>
<th>Mean age in years (range)</th>
<th>No. of positive cases</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>NFB microscopy</td>
</tr>
<tr>
<td>Immigrants</td>
<td>1050 (603/447)</td>
<td>29.2 (14–61)</td>
<td>34*</td>
</tr>
<tr>
<td>Endemic mf-negative†</td>
<td>50 (30/20)</td>
<td>31.5 (5–59)</td>
<td>0</td>
</tr>
<tr>
<td>Non-endemic mf-negative</td>
<td>50 (25/25)</td>
<td>25.1 (21–49)</td>
<td>0</td>
</tr>
<tr>
<td>Other parasitic infections‡</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Malaria</td>
<td>50 (23/27)</td>
<td>32.1 (14–54)</td>
<td>0</td>
</tr>
<tr>
<td>Hydatid disease</td>
<td>30 (21/9)</td>
<td>33.4 (23–58)</td>
<td>0</td>
</tr>
<tr>
<td>Amoebic abscess</td>
<td>20 (14/6)</td>
<td>29.2 (21–54)</td>
<td>0</td>
</tr>
<tr>
<td>Schistosomiasis</td>
<td>60 (35/25)</td>
<td>28.6 (26–67)</td>
<td>0</td>
</tr>
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*Two cases had *B. malayi* mf and thus were excluded from further analysis.
†Individuals were from filaria-endemic areas and were negative for mf by NFB microscopy.
‡Groups of migrants in Kuwait who had never been exposed to filarial infection and were negative for mf by microscopy but positive for other parasitic infections.

Table 2. Performance of NFB microscopy, the ICT and the TropBio assay in detecting filarial infection in the migrant population in Kuwait

Sensitivity and specificity were calculated using ML methods and Staquet’s equation as described in Methods.
and technical constraints, night-time blood specimens were only collected from 25 NFB-negative subjects with CFA and retested for mf by microscopy. Only two of these 25 subjects showed mf with a density of $< 50$ mf ml$^{-1}$. More than 90% of our study subjects were from filaria-endemic areas (Andhra Pradesh, Bihar, Tamil Nadu and Uttar Pradesh – the filaria belt) in India. The microfilaria infection rate in our study correlated well with that reported from these areas (World Health Organization, 1999); however, the prevalence of CFA in our study group was much higher than that of 10·6% observed in the Indian expatriate population in Saudi Arabia (Omar et al., 2000). This is probably due to inclusion of groups of individuals from Kerala in the Indian study, where filarial infection is very low. In this study, 162 (15·9%) mf-negative individuals showed the presence of CFA using the ICT. However, the ICT failed to detect two of the 32 mf-positive cases. The microfilaria load in both of these cases was $< 20$ mf ml$^{-1}$, suggesting a decreased sensitivity of the ICT in cases with a low mf density. The low sensitivity of the ICT was reported previously from India in cases with low mf density (Sunish et al., 2001). We observed an overall sensitivity of 93·8% for the ICT.

The estimated value for specificity of the ICT ranged from 84 to 100% based on the prevalence estimated by NFB microscopy and the ICT. Microscopy detects only infections with patent mf circulating during the night. However, asymptomatic microfilaraemic infections have also been well documented (Eigege et al., 2003). Thus, the ‘false positives’ obtained by the ICT could well be true positives due to absence of mf in the daytime blood samples or the presence of adult worms in the deep tissues. Similarly, the TropBio assay, which detects a different CFA, also detected a much higher prevalence of CFA than mf (20·3 vs 3%, respectively). Earlier studies have also reported similar results (Weerasaooriya et al., 2003). There are several possible explanations for the mf-negative status of these infections. Some infections may be at the pre-patent stage and may be a single-sex infection. Another possibility is that a low level of adult worm infection produces antigenaemia but no detectable microfilaraemia (Eigege et al., 2003; Molyneux & Taylor, 2001). Furthermore, several studies have reported that a considerable number of individuals remain antigen-positive for up to 2 years after diethylcarbamazine treatment (Eigege et al., 2003; Hoerauf, 2003; Meyrowitsch et al., 2004). However, recently one study reported a significant decrease in CFA levels at 18 months after diethylcarbamazine treatment among Myanmar migrants in Thailand (Koyadun et al., 2003).

The performance of the ICT and the TropBio assay to detect CFA showed a statistically significant correlation (192 cases vs 213 cases, respectively); a similar observation has been reported previously (Molyneux & Taylor, 2001).

Although the TropBio assay provides a semi-quantitative response, probably reflecting the adult worm load, it requires more elaborate laboratory facilities and a well-equipped technical staff. The ICT requires minimal laboratory and technical support and thus is more suitable for diagnosis of filarial infection in clinical laboratories and for surveillance studies. In conclusion, this study demonstrated the high specificity and sensitivity of the ICT for detection of filarial infection. The ICT should be used to screen migrant workers in Kuwait for filarial infection using daytime blood specimens.

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