Comparison of tests for the detection of circulating filarial antigen (Og4C3-ELISA and AD12-ICT) and ultrasound in diagnosis of lymphatic filariasis in individuals with microfilariae

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Significant advances were made in the diagnosis of filariasis in the 1990s with the emergence of three new alternative tools: ultrasound and tests to detect circulating antigen using two monoclonal antibodies, Og4C3 and AD12-ICT-card. This study aimed to identify which of these methods is the most sensitive for diagnosis of infection. A total of 256 individuals, all male and carrying microfilariae (1-15,679 MF/mL), diagnosed by nocturnal venous blood samples, were tested by all three techniques. The tests for circulating filarial antigen concurred 100% and correctly identified 246/256 (96.69%) of the positive individuals, while ultrasound detected only 186/256 (73.44%). Of the circulating antigen tests, ICT-card was the most convenient method for identification of Wuchereria bancrofti carriers. It was easy to perform, practical and quick.

Key words: Wuchereria bancrofti - circulating filarial antigen - ultrasound - diagnosis

Lymphatic filariasis (LF) is a parasitic disease caused by nematodes (Wuchereria bancrofti, Brugia malayi and Brugia timori) whose preferred habitats are the lymphatic vessels and lymph nodes which induce the development of disfiguring and debilitating clinical symptoms. The infection, which is transmitted by various genera of mosquitoes (Sasa 1976), is considered to be one of the health problems of greatest social and economic impact in endemic areas (Ottesen et al. 1997).

Until the 1980s, the only direct way to confirm a diagnosis of infection by W. bancrofti was via the identification of microfilariae (MF) in peripheral blood using camera counting, polycarbonate membrane filtration, the thick smear method or Knott’s technique (Knott 1939, Denham et al. 1971, Dennis & Kaen 1971, Eberhard & Lammie 1991). Apart from the low sensitivities of these methods arising from the fact that they only identify filarial infection by way of microfilaraemia, they are also inconvenient in terms of the time of day the blood must be collected, namely when the MF are at a peak in peripheral blood, a parameter that varies from one endemic area to another. In Brazil, the peak for microfilaraemia occurs between 11 pm-1 am (Dreyer et al. 1996).

In the 1990s, significant advances were made in the diagnosis of LF with the emergence of new diagnostic tools: (i) use of recombinant antigens to detect specific antibodies (Chandrasherkar et al. 1994); (ii) a PCR for the detection of filarial DNA (Zhong et al. 1996, Rocha et al. 2002); (iii) the visualisation of live adult worms (AW) using ultrasound (US) and (iv) circulating filarial antigen (CFA) detection (More & Copeman 1990, Weil et al. 1997). At present, the standard diagnostic tools are US (Amaral et al. 1994) and CFA detection, with the latter using the monoclonal antibodies (McAbs) Og4C3 and AD12. The CFA detection techniques are commercially available in the form of kits and have the advantage of allowing for diagnosis to be carried out using blood samples collected at any time of day (More & Copeman 1990, Amaral et al. 1994, Weil et al. 1997, Rocha 2002, 2004).

Og4C3, the first commercially available laboratory kit, is a serological test based on the detection of CFA using an immunoenzymatic technique (ELISA) (More & Copeman 1990) and a McAb of the IgM class of immunoglobulins produced against antigens of the bovine parasite Onchocerca gibsoni. This technique allows for the identification of W. bancrofti antigens in serum, plasma and hydrocele fluid and does not show any cross-reactivity with other helminthic infections (More & Copeman 1990, Rocha et al. 1996, TropBIO 1996, Rocha 2004).

The other serological test, the immunochromatographic or ICT card test, uses the McAb AD12, which recognises a 200 kDa filarial antigen (Weil et al. 1997). The technique is easy to perform, requires no equipment and was launched by ICT Diagnostic (Balgowlah, New South Wales, Australia). It is currently produced by BINAX as NOW® Filariasis in the United States of America.

Another tool for diagnosis of filariasis infection, first discovered by Brazilian researchers, is the use of US to visualise live AW of W. bancrofti in the lymphatic vessels (Amaral et al. 1994, Norões et al. 1996).
Although several authors attest to the high sensitivity of these new methods for diagnosing filarial infection by *W. bancrofti*, a comparative analysis of the sensitivity of these techniques has not been performed.

**PATIENTS, MATERIALS AND METHODS**

*Area and population studied* - This study was carried out on 256 male patients with microfilaremia who were residents of endemic areas in the Metropolitan Region of Recife and referred to the National Center of Lymphatic Filariasis (NCLF)'s outpatient clinic at the Aggeu Magalhães-Research Centre Oswaldo Cruz Foundation (Fiocruz). The patients selected for this study were seen between 2001-2004.

*Parasitological blood test* - After participants had signed the terms of consent, a sample of 5 mL of venous blood was collected between 11 pm-1 am in order to test for the presence of circulating MF. A quantity of 1 mL of blood was filtered through a polycarbonate membrane 13 mL in diameter with 3-µm pores, as described by Rocha et al. (2004).

*Test for live AW* - All of the 256 individuals underwent an US of the scrotal area using a 7.5 MHz probe to visualise the live AW and identify their characteristic movements in lymphatic vessels, i.e., the filariae dance sign (FDS) (Amaral et al. 1994, Norões et al. 1996).

*Collection of serum samples* - One day after venous blood was collected for detection of the presence of MF, an additional sample of 5 mL of venous blood was collected and centrifuged at 2,500 rpm at 20°C; the serum was separated, quantified, labelled and stored at -20°C in the NCLF’s biological specimen bank. In order to detect filarial antigens, the serum from each individual was thawed in an ice bath and the technical procedures for the CFA tests were carried out and processed using a single technique. In comparison, a single serum sample from each individual was exposed to both CFA tests.

*The Og4C3-ELISA test* - The technical procedures were carried out as recommended by the manufacturers of the kit (TropBio®, JCU Tropical Biotechnology Pty Ltd, Townsville, Queensland, Australia). Serological samples were processed in duplicate and the results were obtained using optical density (OD) measurement. The OD measurements for each sample were used to determine the response in units of antigen per mL (ag/mL) by comparison to a standard concentration curve for *O. gibsoni* ag/mL, included in the kit. In accordance with the TropBio norms (1996) samples with ag/mL ≥ 128 were considered positive for CFA.

*The AD12-ICT card test* - The NOW® Filarisis version of the card test (ICT filariasis for blood, serum or plasma, cat. n° 620-000) was carried out according to the manufacturer’s instructions. In short, 100 µL of serum was placed in the position recommended for the test. After one minute, the card was closed and the result was read exactly 10 min later. The test was considered positive when both lines (test and control) could be read through the visualisation window. Any line (light or dark) appearing in the test position indicates that the result of the test is positive; it is negative when the control line can be seen.

*Statistical analysis* - The difference between the proportions was verified using the chi-squared test and the p value at a level of α = 0.05. Owing to the large variation in values for the units of circulating antigen per mL, the differences between the means were tested using the Kruskall-Wallis test.

*Ethical considerations* - This study was approved by the Aggeu Magalhães-Fiocruz Research Centre’s Ethical Committee. The participants (or their parents in the case of minors) were given information about the research and were asked to read and sign the terms of consent. All participants were treated with diethylcarbamazine (6 mg/kg/day/12 days).

**RESULTS**

The age of individuals varied from 7-78 years (mean = 24.14 ± 10.13). The density of MF per mL of blood was between 1-15,679 (mean = 768.56 ± 1.520.24) and the antigenemia levels by Og4C3 varied between 1-56.411 ag/mL (mean = 28.328.87 ± 14.932.78). The percentages for positive results for the three tests are shown in Table I. The sensitivities of the two techniques for detection of CFA (Og4C3 and ICT card) were similar, both showing a high degree of sensitivity (96.69%). The percentage of US positive results was 73.44% (χ² = 4.21; p = 0.046). Live AW was not detected in 68 (26.56%) individuals through the US exam.

The relation between the density of MF/mL and the quantitative and qualitative results of the Og4C3 tests (unit of ag/mL) and the ICT card (positive/negative), respectively, showed a significant increase in the antigen level in Og4C3 CFA and in the percentage of positive tests using the ICT card proportional to the rise in the parasite load chiefly when the microfilaremia density was greater than 100 MF/mL of blood (Tables II, III).

The comparison between US and the density of MF per mL showed that the percentage of tests in which the FDS could be seen significantly increased in proportion to the number of MF. In the patients with a microfilarial density of > 500 MF/mL, AW could be visualised by US in 82.56% (Table IV).

| TABLE I |
| Percentage of positive results and confidence interval (CI) for the ICT card, Og4C3 and ultrasound tests among MF+ mean |

<table>
<thead>
<tr>
<th>Test</th>
<th>(n = 256)</th>
<th>Positives</th>
<th>%</th>
<th>CI 95%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Og4C3-ELISA</td>
<td>246</td>
<td>96.69</td>
<td>93.15-97.99</td>
<td></td>
</tr>
<tr>
<td>ICT card test</td>
<td>246</td>
<td>96.69</td>
<td>93.16-97.99</td>
<td></td>
</tr>
<tr>
<td>Ultrasound</td>
<td>188</td>
<td>73.44</td>
<td>67.77-78.57</td>
<td></td>
</tr>
</tbody>
</table>

a: individuals with microfilaremia diagnosed using the polycarbonate membrane filtration technique; χ² = 4.21; p = 0.046.
A higher proportion of positive results by US was observed in individuals between 20-30 (132/157) and > 30 years (21/35) of age compared to the group between 7-19 years (35/64) of age ($\chi^2 = 23.88$, $p < 0.0001$). In the CFA test results (Og4C3 and ICT card), no statistically significant difference in the proportion of positive results among these age groups was observed ($\chi^2 = 0.63$, $p < 0.731$).

### TABLE II

Average of Og4C3 antigenemia levels according to density of microfilariae

<table>
<thead>
<tr>
<th>Density Microfilariae/mL</th>
<th>n</th>
<th>Mean*</th>
<th>Min</th>
<th>Max</th>
<th>Standard deviation</th>
</tr>
</thead>
<tbody>
<tr>
<td>1-99</td>
<td>78</td>
<td>20,517</td>
<td>1</td>
<td>50,485</td>
<td>15,136</td>
</tr>
<tr>
<td>100-500</td>
<td>82</td>
<td>29,089</td>
<td>3</td>
<td>53,333</td>
<td>14,707</td>
</tr>
<tr>
<td>&gt; 500</td>
<td>86</td>
<td>35,497</td>
<td>1,415</td>
<td>56,411</td>
<td>10,546</td>
</tr>
<tr>
<td>Total</td>
<td>246</td>
<td>28,328</td>
<td>1</td>
<td>56,411</td>
<td>14,933</td>
</tr>
</tbody>
</table>

*a*: arithmetic mean. Kruskal Wallis (2 gL) = 52.00; $p = 0.0001.$

### TABLE III

Percentage of positive results using the ICT card according to the parasite load of microfilariae among individuals carrying microfilariae

<table>
<thead>
<tr>
<th>Density Microfilariae/mL</th>
<th>Positives</th>
<th>%</th>
<th>Negatives</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>1-99</td>
<td>78</td>
<td>89.66</td>
<td>9</td>
<td>87</td>
</tr>
<tr>
<td>100-500</td>
<td>82</td>
<td>98.80</td>
<td>1</td>
<td>83</td>
</tr>
<tr>
<td>&gt; 500</td>
<td>86</td>
<td>100.0</td>
<td>0</td>
<td>87</td>
</tr>
<tr>
<td>Total</td>
<td>246</td>
<td>96.67</td>
<td>10</td>
<td>256</td>
</tr>
</tbody>
</table>

$\chi^2$ test for trend = 12.32; $p = 0.0004.$

### TABLE IV

Frequency of filariae dance sign detected by ultrasound of the scrotum according to microfilariae parasite load in individuals carrying microfilariae

<table>
<thead>
<tr>
<th>Density Microfilariae/mL</th>
<th>Ultrasound (live adult worm)</th>
<th>Positive</th>
<th>%</th>
<th>Negative</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>1-99</td>
<td>55</td>
<td>63.22</td>
<td>32</td>
<td>87</td>
<td></td>
</tr>
<tr>
<td>100-500</td>
<td>62</td>
<td>74.70</td>
<td>21</td>
<td>83</td>
<td></td>
</tr>
<tr>
<td>&gt; 500</td>
<td>71</td>
<td>82.56</td>
<td>15</td>
<td>86</td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>188</td>
<td>73.44</td>
<td>68</td>
<td>256</td>
<td></td>
</tr>
</tbody>
</table>

$\chi^2$ test for trend = 8.266; $p = 0.0040.$

## DISCUSSION

Recent advances in various areas of knowledge related LF have led to the use of new approaches to lessen the severity and impact of the disease (Ottesen et al. 1997). The introduction of new diagnostic tools for detection of infection by *W. bancrofti* in the human population have opened up new possibilities in terms of interrupting transmission and consequently the elimination of filariasis (WHO 1994, Cox 2000, Molyneux et al. 2000, Ottesen 2000). This was reaffirmed in 1997 by the World Health Organisation (WHO), when it established the goal of filariasis elimination worldwide by 2020 (Ottesen 2000).

Until the early 1990s, the only available parasitological method for the evaluation of individuals or populations infected with *W. bancrofti* was the blood thick smear, which uses around 60 µL of capillary blood (Eberhard & Lammie 1991). However, due to its low sensitivity, this method does not allow accurate determination of the true prevalence of infection (McCarthy 2000). With the introduction of the filtration technique and the use of venous blood (up to 10 mL) collected between 11 pm-1 am, a higher degree of reliability of the parasitological status of individuals was obtained (Rocha 2004).

The introduction of US in Recife, Brazil, as a new parasitological tool for diagnosis of bancroftosis made it possible to identify individuals with live AW of *W. bancrofti* in the lymphatic vessels of the scrotal sac (Amaral et al. 1994). Several studies have emphasised the importance of US as a non-invasive parasitological tool (Dreyer et al. 1995, 1998, Faris et al. 1998). The data presented here showed that the use of US was capable of identifying AW in 73.44% of MF+ individuals. This result is similar to that of Norões et al. 1996, who identified AW in 80% of MF+ individuals using US.

The CFA method was more sensitive than the US method in identifying infected individuals. This result is possibly due to the fact that the AW could be located in sites of the body other than male genitals, such as the scrotum, which has been described as the preferred location of AW (Amaral et al. 1994).

Due to the simplicity and convenience of this method, it is possible to carry out the test using blood collected at any time of day. CFA is now considered to be a promising tool for diagnosis of bancroftosis at both the individual and community levels. Among CFA tests, the ICT card, because of its simplicity and precision, has been adopted as an important tool for community surveys and rapid assessment of filarial endemicity in bancroftian filariasis control programs (Ottesen 2000, Molyneux & Taylor 2001). Nevertheless, in large surveys, it can be difficult to examine the cards while at the same time handling many patients, resulting in a tendency of health care providers to gather the cards to be read at a later time. This procedure could contribute to misinterpretation of the results and could increase the number of false positives. In our study, the ICT card test failed to detect CFA in nine of 78 individuals with MF count less than 100 MF/mL, while US failed to detect live AW in 32 of 87 individuals carrying MF. There is concern that currently available tools may have severe limitations when parasite prevalence and antigen levels are low.
Og4C3 and AD12 McAbs detect circulating *W. bancrofti* antigens originating particularly from AW, according to More and Copeman (1990) and Weil et al. (1997). Furthermore, Eberhard et al. (1997) and Lalitha et al. (1997) suggest that the MF do not contribute to the concentration of CFA and that the decrease in the quantity of CFA occurs exclusively as a result of the death of AW. Wong and Guest (1969) noted that microfilarial antibodies were present in all patients with elephantiasis, but were found in none of the patients with circulating MF. Smithers (1968) has suggested that living MF are relatively inert antigenically and do not stimulate the production of antibodies. Our results contradict these observations and clearly show that there is a significant association (p = 0.001) between the density of MF and the concentration of CFA. Similar results were obtained by Rocha et al. (1996), Nicolas et al. (1997) and Itoh et al. (1999). We also agree with Grove and Davis (1978), who suggest that the detection of microfilarial antibodies depends upon the relative release of MF into the bloodstream and the production of antibodies against those MF.

Medeiros et al. (1999) showed that men between 20-30 years of age are more susceptible to LF than are women of the same age group. This was also the largest age group in our study (~ 60%).

The sensitivity of US was lower (73.44%) than that of the monoclonal Og4C3 (96.69%) and AD12 (96.69%) techniques. On the other hand, for age groups 20-30 and > 30 years, the US was statistically significantly (p > 0.0001) effective in detecting AW. Therefore, in order to provide a rapid and reliable detection of infection with *W. bancrofti*, the choice of diagnostic tool must take into consideration age-group, gender, the presence of infected individuals harbouring low levels of MF, the presence of growing worms or non-fecund adults or adults of a single sex and the endemicity of the area.

In summary, in our study, McAbs Og4C3 and AD12 showed high sensitivity for detecting infected individuals. While the global LF elimination program is ongoing, highly sensitive and specific diagnostic tools are necessary to monitor and control the programs. There is presently no “gold standard technique” that alone is capable of offering complete confidence that infection has been eliminated and that can by its use alone allow the WHO to certify an area free of LF. The Og4C3 ELISA and the ICT card test, in our study, presented high performance in detecting positive individuals. Nevertheless, both have advantages and inconveniences with respect to their use in large control programs. The ICT card test may be most appropriate, because of its ease of use and because results are obtained within 10 min. In cases where one needs to broaden the diagnostic investigation, US should be of great value, as demonstrated here.

Until now, no CFA test available has been reported to be 100% specific in detecting infection by *W. bancrofti* (Rocha et al. 1996).

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