Diagnosis of toxoplasma infection in cardiac transplant recipients using the polymerase chain reaction

R Holliman, J Johnson, D Savva, N Cary, T Wreghitt

Abstract
Cardiac biopsy samples taken from transplant recipients around the time of primary toxoplasma infection were investigated by conventional histology and amplification of the P30 gene of Toxoplasma gondii by the polymerase chain reaction (PCR). Toxoplasma was detected more frequently by PCR than histology which may reflect the enhanced sensitivity of the former technique. Further studies are required to determine the optimal amount of tissue which should be examined by each technique and to develop a PCR assay capable of distinguishing between quiescent infection and active toxoplasmosis.

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Toxoplasmosis can be life threatening to cardiac transplant recipients. Those at greatest risk are Toxoplasma specific antibody negative recipients who are exposed to the parasite for the first time when receiving viable cysts in the organ graft of seropositive donors. About 60% of such mismatches result in severe toxoplasma infection, although this incidence is reduced significantly when prophylaxis with pyrimethamine is given. Primary toxoplasma infection originating in the transplanted heart can be diagnosed by serological or histological methods. The infected recipient usually has detectable specific IgM and rising titres of IgG, but discordant findings have been reported. Histological examination may give false negative results as organisms are usually confined to a few cysts, even in severe, acute infection. The differentiation of toxoplasma infection and acute rejection can be problematic and the two conditions may be concomitant. We used the polymerase chain reaction (PCR) to investigate cardiac biopsy samples in cases of acute toxoplasmosis associated with heart transplantation and compared the findings with those obtained with conventional histological examination.

Methods
Toxoplasma DNA was amplified using the PCR and two sets of oligonucleotide primers based on the published sequence of the P30 gene. Amplification reactions were performed, as described previously, using serial cycles of PCR resulting in a 914 and then a 522 base pair fragment of DNA. Amplified fragments were visualised in ethidium bromide stained gels and by DNA-DNA hybridisation using 32P radiolabelled 522 base pair fragment as a specific probe. Primary toxoplasma infection was diagnosed by serological investigation of toxoplasma specific IgG and IgM. Cardiac biopsy samples were taken from heart transplant recipients around the time of infection and examined by PCR and conventional histological techniques.

Results
A total of 15 biopsy samples taken from seven cardiac transplant recipients with acute toxoplasmosis were investigated. A comparison of histological findings and detection of toxoplasma by PCR is presented in the table.

Comparison of histology and PCR results with serological findings for seven patients with primary donor acquired T gondii infection

<table>
<thead>
<tr>
<th>Case No</th>
<th>Biopsy date</th>
<th>Histological findings</th>
<th>PCR</th>
<th>Timing of T gondii seroconversion (transplantation date)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>02/05/84</td>
<td>Negative</td>
<td>-</td>
<td>22 May 1984</td>
</tr>
<tr>
<td>2</td>
<td>23/05/84</td>
<td>Toxo seen</td>
<td>+</td>
<td>(26/04/84)</td>
</tr>
<tr>
<td></td>
<td>27/04/82</td>
<td>Negative</td>
<td>+</td>
<td>End May 1982</td>
</tr>
<tr>
<td></td>
<td>06/05/82</td>
<td>Negative</td>
<td>+</td>
<td>(20/04/82)</td>
</tr>
<tr>
<td>3</td>
<td>17/05/82</td>
<td>Negative</td>
<td>+</td>
<td>1985-88 Serum samples not available between these dates</td>
</tr>
<tr>
<td></td>
<td>28/07/85</td>
<td>Negative</td>
<td>+</td>
<td>probably took place in 1985 (1807/85)</td>
</tr>
<tr>
<td></td>
<td>09/08/85</td>
<td>Negative</td>
<td>+</td>
<td>March-November 1987</td>
</tr>
<tr>
<td></td>
<td>05/09/85</td>
<td>Negative</td>
<td>-</td>
<td>Serum samples not available between these dates</td>
</tr>
<tr>
<td>4</td>
<td>13/11/87</td>
<td>-</td>
<td>+</td>
<td>(03/09/87)</td>
</tr>
<tr>
<td>5</td>
<td>17/11/87</td>
<td>Negative</td>
<td>-</td>
<td>December 1987-January 1988</td>
</tr>
<tr>
<td>6</td>
<td>12/01/88</td>
<td>Negative</td>
<td>+</td>
<td>(19/09/87)</td>
</tr>
<tr>
<td></td>
<td>21/09/81</td>
<td>Negative</td>
<td>+</td>
<td>End October 1991</td>
</tr>
<tr>
<td></td>
<td>29/09/81</td>
<td>Toxo seen</td>
<td>+</td>
<td>(10/09/81)</td>
</tr>
<tr>
<td>7</td>
<td>30/06/83</td>
<td>Negative</td>
<td>-</td>
<td>Mid July 1983</td>
</tr>
<tr>
<td></td>
<td>26/07/83</td>
<td>Negative</td>
<td>-</td>
<td>(22/06/83)</td>
</tr>
</tbody>
</table>
Toxoplasma was detected in two of the seven cases by histology and in five cases using PCR.

Conclusions
Previous studies have shown that a single toxoplasma organism can be detected by PCR and that the method is highly specific. Contamination of the reactions leading to false positive results was unlikely as control samples of known toxoplasma free tissue repeatedly failed to produce amplification products. Consequently the enhanced detection of toxoplasma by PCR compared with histology may reflect the superior sensitivity of the former technique. In its present form the PCR assay will not distinguish quiescent cysts from active infection with free trophozoites of *T. gondii*. This problem may be overcome by the development of stage specific primers. As the distribution of toxoplasma in infected tissue is uneven, sampling error could result in discordant histological and PCR findings. The optimal amount of tissue which should be examined by each technique is not established.

Our results indicate that PCR can be used to detect *T. gondii* in cardiac biopsy samples taken from transplant recipients with primary, organ graft related toxoplasmosis. This method may be more sensitive compared with conventional histological examination. Further studies are in hand to define the application of PCR in organ transplantation for the diagnosis of primary infection and the identification of organ grafts which contain the parasite.

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**Specificity of colon specific antigen and colon ovarian tumour antigen**

A M Dorman, D Chin, M Leader

Abstract
Sections (5 μm thick) from 101 primary adenocarcinomas (including ovarian, colorectal, gastric, breast, oesophageal, prostatic, pancreatic, endometrial and gall bladder) were incubated with anti-colon specific antigen (CSA) and anti-colon ovarian tumour antigen (COTA) antibodies using the peroxidase antiperoxidase technique with positive and negative controls. Anti-CSA positivity was seen in 19 of 20 colonic adenocarcinomas, but it was also seen in a large number of the other tumours. While anti-COTA staining was positive in 16 of 20 colonic adenocarcinomas and 20 of 30 ovarian adenocarcinomas, it was also positive in a large number of the tumours.

Anti-CSA and anti-COTA are not adequately specific in the identification of a colonic or ovarian origin of an adenocarcinoma and cannot reliably be applied to the identification of a metastatic adenocarcinoma of unknown primary site.

A patient with metastatic adenocarcinoma of unknown primary site is a relatively common clinical problem. The identification of an antigen specific for a primary site, such as colon or ovary would have the benefit of minimising invasive and expensive investigations in untreatable patients and would also indicate the surgical and chemotherapeutic regimens in potentially curable patients.

Since the discovery of carcinoembryonic antigen (CEA) in colon cancer other more allegedly specific colon cancer associated antigens (colon specific antigen (CSA) and colon ovarian tumour antigen (COTA)) have been identified.

CSA is a heat stable, mucin associated antigen present in normal colonic epithelial cells and is expressed in greater quantities in colonic adenocarcinomas. COTA is a heat stable antigen present in colonic adenomatous polyps and adenocarcinomas but not in normal colonic epithelium and it is also present in mucinous ovarian tumours.

In 1986 Pant *et al* described the sensitivity and specificity of goat anti-COTA antibody to
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