Anti-sperm antibodies in homosexual men: prevalence and correlation with sexual behaviour

B P Mulhall, S Fieldhouse, S Clark, L Carter, L Harrison, B Donovan, R V Short

Abstract

The sera of 60 homosexual males were examined for the presence of antibodies to sperm using an indirect immunobead test (IBT). Six of 60 (10%) had antibodies of IgG isotype; in addition two of the six had antibodies of IgA isotype. The presence of antibodies was associated with the practice of unprotected receptive anal intercourse in the previous six months. Antibodies were not found in homosexual men who were celibate, or who practised only oral intercourse during the same period. There was no correlation between the presence of anti-sperm antibodies and antibodies to human immunodeficiency virus (HIV), or numbers of T lymphocytes. These preliminary results lend support to the hypothesis that antigen presentation in the lower gut may be a source of sensitisation against sperm. The possibility that anti-sperm antibodies may be a marker of receptive anal intercourse merits further investigation.

Antibodies to various components of sperm and seminal plasma are present in the sera of certain individuals, particularly vasectomised men and infertile couples, and there has been much speculation over their pathogenetic role. Recently, using a variety of methods, anti-sperm antibodies have also been detected in the sera of healthy homosexual men, or those who presented with HIV infection, or (other) sexually transmitted diseases. Together with data from experimental animal models, therefore, it has been suggested that the introduction of spermatozoa into the rectum may lead to the development of a humoral immune response. However, many of these earlier studies either lacked information on the subjects' HIV status or used methods for measuring sperm antibodies that were only poorly reproducible, and with the exception of one study did not attempt to correlate the findings with specific sexual practices. A recent editorial stated that the most useful assay for anti-sperm antibodies is the immunobead method and this prompted us to use this technique to investigate sperm antibody titre and prevalence in a group of well documented homosexual men, to see to what extent it is correlated with recent sexual behaviour, HIV status, and indices of immune function.

Patients

Between January and March 1988, sixty four homosexuals (range 28–60 yr, mean 40·7) attending the Royal Melbourne Hospital HIV clinic as part of an ongoing epidemiological study completed questionnaires on their sexual practices over the previous six months. Based on their responses men were categorised into the following groups: Group 1—unprotected oral intercourse only (n = 13), Group 2—unprotected oral and anal intercourse (n = 39), Group 3—unprotected anal intercourse only (n = 2), and Group 4—celibate (n = 6).

All individuals in the first three groups were recipients of sperm; six men who reported always using condoms during the previous six months were excluded from the study. On the basis of informed consent, blood was obtained for anti-sperm and anti-HIV antibody testing as well as for quantitation of lymphocyte subsets. In addition sera were obtained from two homosexual men who had attended the Melbourne Sexually Transmissible Diseases Centre for treatment of rectal gonorrhoea; these were placed in Group 2.

Methods

Fresh donor sperm was obtained from volunteers. These were HIV antibody negative heterosexual men who were regular sperm donors in an in-vitro fertilisation program. Test sera were screened for anti-sperm antibodies using a minor modification of the indirect immunobead assay previously described.
This is essentially a rosetting technique: briefly, serum at a dilution of 1:10 was incubated for 30 min at 37°C with live sperm (count > 60 x 10⁶/ml and motility > 50%) from donors who were anti-sperm antibody negative (determined by incubation of sperm with IgG or IgA coated immunobeads in the absence of serum). After washing to remove free immunoglobulin, the sperm were added to immunobeads coated with rabbit anti-human IgG or IgA (Biorad, Ca), and under magnification (× 25), 100 motile sperm were counted. The result was considered positive if > 10% of motile sperm had two or more beads attached. In each assay positive and negative control sera were included. Positive control sera were obtained from heterosexual men attending an infertility clinic. To minimise interassay variation results were expressed relative to a positive control value of 100%. In addition, all positive samples were precipitated with PEG, ammonium sulphate, and more specifically, anti IgG or IgA coated beads, to check that anti-sperm activity resided in the immunoglobulin fraction. All positive sera were also serially diluted to determine anti-sperm antibody titre. Sera were screened for antibodies to HIV using an ELISA assay and confirmed by Western Blot. At the same time absolute numbers of CD4+ and CD8+ T lymphocytes were determined using the monoclonal antibodies Leu 3a and Leu 2 (Becton Dickinson, Ca) and flow cytometry (Coulter Epics IV, Ca).

**Results**

Six of 60 (10%) sera tested positive for antibodies to sperm. All six had antibodies of IgG isotype; two of the six also had antibodies of IgA isotype. For IgG, three sera were positive at a dilution of 1:100 and the remaining three at a dilution of 1:50. The positive control sera were positive at a dilution of 1:500. For IgA, titres were lower in controls (1:200) and in patients (1:50). Anti-sperm activity was completely removed in all cases by precipitation with coated beads. Three different sperm donors were used and the inter-donor variation was less than 5%.

All six subjects positive for anti-sperm antibodies were from Groups 2 and 3 (table 1), viz. they practised receptive anal intercourse; none of the thirteen recipients of oral intercourse alone had antisperm antibodies. Thirty subjects had antibodies to HIV; clinically they were all asymptomatic, that is, they belonged to the Centers for Disease Control (CDC) Groups II and III. Of the six antisperm antibody positive sera, two were positive and four negative for antibodies to HIV (table 2). However, given the small numbers, none of these results attained statistical significance (chi square test with Yates’s correction and Fisher’s exact test). There was a high significant difference in indices of immune function between the HIV seropositive and seronegative groups with respect to absolute numbers of CD4+ (p < 0.01) and CD8+ (p < 0.005) T lymphocytes but not between anti-sperm antibody positive and negative subjects. There were no significant differences in age between any of the groups.

**Table 2 Anti-sperm antibodies and HIV status**

<table>
<thead>
<tr>
<th>HIV status</th>
<th>+ve</th>
<th>−ve</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anti-sperm antibody status</td>
<td>6 2</td>
<td>4 28</td>
</tr>
</tbody>
</table>

Anti-sperm antibodies compared with the presence of antibodies to HIV.

**Discussion**

The prevalence of antibodies to sperm in this group of homosexual men was 10%, and was confined to 17% of those who had practised unprotected anal receptive intercourse in the previous six months. Other investigations using different methods have found prevalence rates of 20–50% in similar groups, although their studies were performed earlier in the HIV epidemic, before the availability of HIV testing, and when subjects were more likely to have practised unprotected anal receptive intercourse more frequently. In addition “naturally occurring” antibodies have been detected in the general male and female population. Previous methods for detecting anti-sperm antibodies have included sperm immobilisation, agglutination, indirect immunofluorescence, enzyme-linked immunosorbent assays (ELISA) and Western blotting, and have been striking in their lack of consistency and correlation, particularly when different target sperm preparations were used. In the present study, employing the immunobead test, the variation between three different sperm donors was less than 5%. However, there is every likelihood that the antibodies recognised in the immunobead test are different from those detected in previous assays. In addition, the prevalence of these antibodies in the heterosexual population has not been fully investigated.

Experimental studies in rabbits have demon-
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Address for reprints: Dr B P Mulhall, Department of Public Health, University of Sydney, NSW 2006, Australia.


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