DETECTION OF HIV AND HCV RNA IN SEMEN FROM BRAZILIAN COINFECTED MEN USING MULTIPLEX PCR BEFORE AND AFTER SEMEN WASHING

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SUMMARY

Introduction. Prolonged survival of patients under HAART has resulted in new demands for assisted reproductive technologies. HIV serodiscordant couples wish to make use of assisted reproduction techniques in order to avoid viral transmission to the partner or to the newborn. It is therefore essential to test the effectiveness of techniques aimed at reducing HIV and HCV loads in infected semen using molecular biology tests.

Methods. After seminal analysis, semen samples from 20 coinfected patients were submitted to cell fractioning and isolation of motile spermatozoa by density gradient centrifugation and swim-up. HIV and HCV RNA detection tests were performed with RNA obtained from sperm, seminal plasma and total semen.

Results. In pre-washing semen, HIV RNA was detected in 100% of total semen samples, whereas HCV RNA was concomitantly amplified in only one specimen. Neither HIV nor HCV were detected either in the swim-up or in the post-washing semen fractions.

Conclusions. Reduction of HIV and/or HCV shedding in semen by density gradient centrifugation followed by swim-up is an efficient method. These findings lead us to believe that, although semen is rarely found to contain HCV, semen processing is highly beneficial for HIV/HCV coinfected individuals.

KEYWORDS: HIV; HCV; Coinfection; Serodiscordant couples; Semen washing; Swim-up.

INTRODUCTION

According to current figures in the Brazilian Epidemiological Reports (www.aids.gov.br), there are 600,000 HIV-infected people in Brazil, of whom 65% are males of sexually active age. Also worthy of note is the growing proportion of Brazilians (17.7% to 36.2%) who are also infected with the hepatitis C virus (HCV). This coinfection is an important factor in the natural history of AIDS and contributes to greater morbidity and mortality in these patients. Furthermore, prolonged survival of patients under HAART (Highly Active Antiretroviral Therapy) has given rise to a new demand from serodiscordant couples who wish to have children. As these couples cannot stop using condoms, lest they infect their seronegative partner or the newborn, it is essential to treat the infected semen with effective laboratory techniques that not only isolate the best spermatozoa, but also remove HIV and HCV infected cells from the semen.

This is achieved in many studies by using the semen washing technique together with the swim up method to purify the infected semen. These techniques are currently in widespread use in different parts of the world. Some authors, however, question this type of procedure and raise doubts as to its effectiveness in removing HIV from sperm. There are still questions regarding the possible passive transport of the virus from the systemic compartment to the genital tract. Several studies report both a limited relationship between blood plasma and seminal-HIV viral load, as well as the presence of phylogenetically different strains in both. They also report possible intermittent excretion of HIV in semen, whereas in blood the viremia persists during all stages of the disease, even during HAART. Nevertheless, a small fraction of HIV carriers remain asymptomatic after years of infection, with a CD4+ cell count of over 500 mm³ and undetectable viral load, even without any treatment. These patients are designated long-term non-progressors (LTNPs).

The risk of hepatitis C transmission in assisted reproduction techniques is currently the subject of widespread debate. LEVY et al. investigated the presence of HCV RNA in the semen of infected men in order to find out if semen fractions act as a reservoir for the hepatitis C virus. In their study, HCV RNA was found in 2/39 (5%) of semen and semen fraction samples, giving direct evidence of the risk of contamination that may exist in human reproduction techniques. Other studies concluded that although the presence of HCV in semen...
is intermittent and not common, sexual transmission cannot be ruled out.

In contrast, both recent and earlier studies concluded that purification of motile spermatozoa by either the discontinuous gradient or swim-up method, or even by a combination of these techniques, reduces the number of HIV and/or HCV particles in semen to undetectable levels, making it possible for serodiscordant couples to conceive a child by means of assisted reproduction techniques with minimum risk of seroconversion.

Although there are a large number of studies regarding the transmission of HIV and/or HCV through assisted reproduction techniques, the findings differ and are to a certain extent in conflict with each other, thus reinforcing the need for further investigation using molecular biology tests.

**OBJECTIVE**

This study set out to evaluate the effectiveness of density gradient centrifugation and swim-up techniques in reducing HIV and HCV loads in infected semen, using molecular biology tests for viral detection.

**MATERIAL AND METHODS**

1. **Population selection:** Between January 2003 and May 2004, semen was collected from a convenience sample of 20 HIV/HCV-coinfected patients in the 27- to 45-year age range who had undetectable plasma viral loads for HIV, were positive for HCV RNA in blood (by RT-PCR) and for anti-HCV (by ELISA), and who had presented at the Human Reproduction Section of the Federal University of Sao Paulo for assisted reproduction screening. Patients who had undergone vasectomy, had azoospermia or referred a symptomatic Sexually Transmitted Disease (STD) at enrolment were excluded from the study. The investigation protocol was approved by the Institutional Review Board.

2. **Semen collection and processing:** Samples were collected by masturbation into a sterile polypropylene flask after three to five days of ejaculatory abstinence. After liquefaction and homogenization, the seminal sample was divided into two aliquots. The first aliquot was initially submitted to semen analysis according to the World Health Organization (WHO) guidelines. Following semen analysis, the aliquot was submitted to semen processing using a discontinuous Percoll gradient technique at two different concentrations (90% and 45%). After centrifugation (110 g, 30 minutes), three different fractions were obtained: seminal plasma, dead sperm and cellular elements other than sperm (germ line cells, cellular debris, bacteria and leukocytes), and motile spermatozoa.

The second aliquot (total semen) was stored in a -70 °C freezer and used later in parallel with seminal fractions in our search for HCV and HIV genital shedding. For this purpose, 100 µL of total semen and of the three fractions obtained by the discontinuous gradient technique were used for RT PCR multiplex amplification of HCV and HIV sequences before semen washing.

The remaining volume of each fraction (approximately 900 µL) was washed three times with RPMI and the cell pellet resuspended in 1 mL of medium. The fraction containing motile spermatozoa was further submitted to the swim-up method. The best motile sperm that migrated to the culture medium were then carefully removed.

All samples (positive control, total semen and seminal fractions) were then submitted to nucleic acid extraction using the Nuclisens kit (Biomerieux, Brazil) according to the manufacturer’s instructions. Nucleic acids were eluted in 50 µL of a specific buffer and stored at -70 °C until qualitative pre- and post-semen-washing RT-PCR viral-sequence amplification was performed. This test was carried out using 100 µL of each pre- and post-semen-washing aliquot.

3. **HIV and HCV Multiplex RT-PCR**

3.1 **cDNA synthesis:** Complementary DNA (cDNA) was synthesized using a reaction mixture that contained 1x RT Buffer, 0.2 mM dNTPs (dATP, dTTP, dCTP, dGTP), 4.0 mM MgCl₂, 40 IU RNAsin, 103M DTT, 2.5 µM, PDN(6) 2.5U, M-MLV RT and 22.0 µL of extracted RNA in a total volume of 40 µL, following a standardized protocol (65 °C for five min, 22 °C for 10 min, 37 °C for 30 min and 95 °C for five min).

3.2 **HIV/HCV Multiplex RT-PCR:** Ten microlitres of cDNA were then submitted to a qualitative multiplex RT-PCR technique, in which three viral genome sequences (HIV, HCV and Dengue type 1 virus) were simultaneously amplified using specific primer pairs for each viral agent.

Multiplex RT-PCR amplification was performed in total semen as well as in the spermatozoa and dead-sperm and nonspermatozoa seminal fractions using a reaction mixture that contained 1x Buffer, 0.12 mM dNTPs (dATP, dTTP, dCTP and dGTP), 2.0 mM MgCl₂, 1.25U of AmpliTaq Gold® (Applied Biosystems, Foster City, CA, USA) in 40 µL total volume. The following amplification protocol was used: one cycle at 95 °C for five minutes followed by 40 cycles consisting of

<table>
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<tr>
<th>Table 1</th>
<th>Specific primer pairs used for multiplex RT-PCR amplification of viral genome sequences in seminal samples</th>
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<tbody>
<tr>
<td>HIV</td>
<td>RAR 1032 5’GAG ACA CCA GGA ATT AGA TAT CAG TAC AAT GT3’</td>
</tr>
<tr>
<td>HIV</td>
<td>RAR 1033 5’CTA AAT CAG ATC CTA CAT ATG CAT CAT GT3’</td>
</tr>
<tr>
<td>HCV</td>
<td>SM3 5’CTA GCC ATG GCG TTA GAT3’</td>
</tr>
<tr>
<td>HCV</td>
<td>HC18 5’GTT GCA CGG TCT ACG AGA CCT3’</td>
</tr>
<tr>
<td>Dengue</td>
<td>D1 5’TCA ATA TG CTA GAC GCG GGA ACC G3’</td>
</tr>
<tr>
<td>Dengue</td>
<td>TS1 5’CGT CTC ATG GAT CCG GGG G3’</td>
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30 seconds at 94 °C, 30 seconds at 55 °C and one minute at 65 °C, and a final 7-minute extension at 65 °C in an automated thermal cycler.

To validate the nucleic-acid extraction and the RT-PCR amplification steps, 10 µL of serum known to be infected by the Dengue I virus were added to each sample as an internal reaction control. We used a 100 µL mix containing 10 µL of HCV-positive (774,892 viral copies/mL) and 10 µL of HIV-positive (1,700,000 viral copies/mL) sera, 10 µL of Dengue I virus culture supernatant and 70 µL of HCV, HIV and dengue-negative serum as a positive control for the HIV/HCV multiplex RT-PCR reaction. Analytic sensitivity is 500 UI/mL for both viruses (José Eduardo Levi, personal communication).

After electrophoresis, RT-PCR amplification products were visualized under ultraviolet light on 2% agarose gels stained with ethidium bromide.

**RESULTS**

Each patient provided one seminal sample that was used for seven different HIV/HCV multiplex RT-PCR tests (total semen in the pre-washing phase; seminal pre and post-washing phases). A total of 140 multiplex RT-PCR reactions were performed on semen samples collected from twenty HIV/HCV-coinfected patients.

In the pre-washing phase HIV RNA was detected in 100% of total-semen samples, as well as in 50% of seminal plasma samples. HIV-RNA sequences were detected in dead-sperm and non-sperm cells from six out of twenty cases (30%) and in motile-spermatozoa fractions in one out of twenty cases (5%). In contrast, HCV RNA was detected in only one of the 20 total-semen specimens tested.

After semen washing, neither HIV RNA nor HCV RNA could be detected in any of seminal fractions. However, Dengue virus sequences were detected in all specimens, confirming the efficiency of the extraction, reverse-transcription and PCR-amplification steps.

**DISCUSSION**

HIV and HCV share the same transmission pathways (parenteral and sexual), and there is therefore an increased risk of coinfection, especially as the hepatitis C virus is ten times more infectious than HIV. HIV/HCV coinfection is a serious public health problem worldwide, and 30% of people infected with HIV are also coinfected with HCV. There is evidence that this coinfection can change the natural history of HIV/AIDS infection by causing the number of TCD4+ blood cells to decrease faster, as HCV leads to proliferation of TCD4+ cells in hepatic tissue and promotes HIV replication. This would especially be the case in patients with high HCV titer, due to interference in the production of cellular cytokines, causing the HIV/HCV viral load to increase even further.
The advent of HAART, however, is also changing the natural history of HIV/HCV coinfection because of increased patient survival. As a result, an ever increasing number of serodiscordant couples wish to have their own child without resorting to gamete donation in sperm banks or giving up safe sex. The only option available to these couples is to seek assisted reproduction techniques associated with ultrasensitive molecular biology tests that ensure that their partner will not be contaminated or that the virus will not be transmitted horizontally or vertically.

Some studies suggest that the presence of HIV is an important cofactor in increased sexual transmission of HCV. The probability of vertical transmission in female coinfected patients is always higher and is always associated with high viral load.

In our study, HCV RNA was detected in the pre-semen washing phase in only one sample of total semen, suggesting that sexual transmission can occur, but that it is less frequent than HIV. These findings agree with data in the literature, where HCV was rarely detected in semen and then only intermittently.

In contrast, there was a 100% prevalence of HIV RNA in total semen. This positivity rate confirms both the high degree of sensitivity of our multiplex PCR and the effectiveness of the HIV/HCV RNA extraction method, which eliminates possible inhibitors that could produce false-negative results.

Because our casuistic only includes patients with undetectable plasma viral load for HIV, the high prevalence of HIV RNA found in total semen can be explained by HIV compartmentalization as and is always associated with high viral load.

We conclude that semen washing, together with the relevant reproductive technology and HAART, reduce the risk of viral transmission. These findings lead us to believe that, although semen is rarely found to contain HCV, semen purification is highly beneficial for HIV/HCV coinfected individuals.

REFERENCES


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