REVIEW

Contraception and HIV: what do we know and what needs to be done?

Laura Waters, Simon Barton

Introduction

The era of highly active antiretroviral therapy (HAART) has associated with dramatic reductions in the rates of morbidity and mortality associated with HIV infection. In countries with access to HAART, the management of HIV has evolved to one where long-term management and related issues form an increasing part of patient care.

In recent years, the greatest number of new HIV diagnoses in the UK have been amongst heterosexuals and, in addition to the huge improvements in the quality of life enjoyed by these individuals, reproductive health forms an increasingly important aspect of patient care. For example, amongst sero-discordant couples (i.e. where only one partner is HIV-infected) the prevention of HIV transmission is paramount when reviewing contraceptive options. Amongst sero-concordant couples (i.e. where both partners are HIV-infected) the potential for transmitting resistant virus must be considered.

The use of HAART is associated with potentially complex interactions with a number of drugs including hormonal contraceptives. In this article we review the contraceptive options available for couples living with HIV infection and, in particular, review the current knowledge on interactions between HAART and hormonal methods of contraception.

HIV and fertility

Whether or not HIV infection affects fertility is uncertain, and studies of female endocrine function have yielded conflicting results. One retrospective study comparing 248 HIV-infected women with 82 controls found irregular bleeding and amenorrhoea were more frequent in the HIV-infected subjects. HIV-related factors such as stage of disease and CD4 cell count were not associated with menstrual irregularities.

In contrast, a retrospective analysis of progesterone and follicle-stimulating hormone levels in stored blood samples estimated ovulation to be more regular in women with higher CD4 cell counts. Finally, Harlow et al. followed 802 HIV-positive and 273 HIV-negative females and studies of female endocrine function have yielded conflicting results. One retrospective study comparing 248 HIV-infected women with 82 controls found irregular bleeding and amenorrhoea were more frequent in the HIV-infected subjects. HIV-related factors such as stage of disease and CD4 cell count were not associated with menstrual irregularities.

In terms of contraceptive efficacy, perfect use of male condoms is associated with a pregnancy risk of 3% per year and ‘real-life’ use with an annual pregnancy rate in the region of 12–14%. Studies reveal breakage rates of up to 6.7% and slippage rates between 0.6% and 13.1%.

Female condoms

These are made of polyurethane and comprise two flexible rings; one fits over the cervix and the other over the vaginal entrance. The cumulative exposure of vaginal mucosa to sperm has been estimated to be lower with female than male condoms (i.e. approximately 3% vs 11.6%).

In terms of contraception, the female condom has been shown to provide similar protection against pregnancy as other barrier methods. One study suggested a higher rate of pregnancy when compared with the male condom, with 12-month failure rates of 5% and 3%, respectively. A recent Japanese study demonstrated a 1% failure rate at...
6 months with consistent use, increasing to 3% with typical use. This equates with an estimated Pearl index of 1.0–5.0.

Other barrier methods
These include the diaphragm, which covers the cervix and parts of the upper vagina, and the cap, which sits over the cervix only. These methods cannot be recommended for HIV-infected individuals due to the large areas of vaginal mucosa that remain exposed. Additionally, the use of spermicides in conjunction with diaphragms or caps may actually increase the risk of HIV transmission.

Nonoxynol-9, a product commonly used in spermicides, was compared with non-spermicidal gel lubricant in a trial of over 750 HIV-negative commercial sex workers. All were supplied with condoms and carefully counselled about their use. By the end of the study period the overall rates of HIV infection were around 50% greater in the nonoxynol-9 group. HIV acquisition was particularly high in the women who had used the spermicide without condoms. The women in the spermicide arm of the study were also found to have more vaginal lesions on examination. This confirms the findings of earlier work by Kreiss et al. who demonstrated a trend for increased HIV acquisition amongst women using nonoxynol-9-impregnated sponge secondary to associated vaginal ulceration.

Despite the clear evidence of increased female HIV acquisition with the use of nonoxynol-9 there is a lack of reciprocal evidence in terms of female-to-male transmission. It is prudent to advise HIV-positive women in a relationship with an HIV-negative partner to avoid the use of nonoxynol-9. Whether this advice extends to spermicidally lubricated condoms is, at present, uncertain.

Hormonal contraception
Hormonal contraception, in the form of the contraceptive pill, is the most frequently used method of contraception in the UK, being used by 26% of women aged between 16 and 49 years. It must be emphasised that all hormonal methods may be affected by drug–drug interactions with anti-retrovirals and do not confer protection against acquisition of sexually transmitted infections (STIs) including HIV.

Combined oral contraceptive pill (COCP)
This is a highly effective method of contraception when used correctly and women with HIV infection fall into World Health Organization (WHO) Category 1, which means they are eligible for unrestricted use. Women using liver enzyme-inducing drugs, however, which includes some anti-retrovirals, become WHO Category 3 (risks outweigh benefits). This classification is applied on the basis of drug interactions as opposed to medical risk.

The ‘perfect use’ failure rate of the COCP is 0.1% and the ‘typical use’ rate is around 5% per year giving an estimated Pearl index of 0.3–4.0 per 100 woman-years. This method may be associated with a decreased risk of osteoporosis and related fractures, since HIV infection per se and possibly some anti-retrovirals, are associated with reductions in bone mineral density, the COCP could confer a degree of protection to HIV-infected women.

Ethinylestradiol (the oestrogenic component in COCP) and progestogens (the exact formulation varies from one brand to another) are metabolised by the CYP 3A4 isoenzyme of the hepatic cytochrome system. Any agent, including many anti-retrovirals, that induces or inhibits this enzyme can therefore decrease or increase hormone levels. Unfortunately these interactions can be unpredictable, especially for agents such as efavirenz that act as inhibitors and inducers, in addition to being substrates of the enzyme themselves.

The current pharmacokinetic evidence relating to anti-retrovirals and hormonal methods of contraception is summarised in Table 1.

Most of these studies are short-term. Longer-term interactions and the clinical significance of the pharmacokinetic changes are unknown. One study demonstrated small reductions in levels of ethinylestradiol levels with concomitant ritonavir (high dose) when 50 µg oestriadiol was used, but no documented pregnancies. In summary, the pharmacokinetic evidence base is limited and

Table 1 Summary of current pharmacokinetic evidence relating to anti-retrovirals and hormonal methods of contraception

<table>
<thead>
<tr>
<th>Anti-retroviral</th>
<th>Interaction with ethinylestradiol</th>
<th>Interaction with progestogens</th>
<th>Advice</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amprenavir (APV)</td>
<td>Reduced APV levels (with Ortho-Novum®); reduced oestradiol</td>
<td>Reduced APV levels; increased norethisterone levels</td>
<td>Avoid COCP as reduced APV levels</td>
</tr>
<tr>
<td>Atazanavir (ATV)</td>
<td>Increased oestradiol levels with unboosted ATV; ATV usually boosted with 100 mg RTV once daily</td>
<td>Increased norethisterone levels with unboosted ATV</td>
<td>No impact on contraceptive efficacy if unboosted; boosted not studied (higher levels of RTV reduce hormone levels)</td>
</tr>
<tr>
<td>Kaletra (lopinavir + ritonavir)</td>
<td>Significantly reduced oestradiol levels</td>
<td>Reduced norethisterone levels</td>
<td>Alternative methods advised</td>
</tr>
<tr>
<td>Nelfinavir (NFV)</td>
<td>Significantly reduced oestradiol levels</td>
<td>Reduced norethisterone levels</td>
<td>Alternative methods advised</td>
</tr>
<tr>
<td>Ritonavir (RTV)</td>
<td>Reduced oestradiol levels with 500 mg RTV twice daily</td>
<td>Nil documented</td>
<td>Advise COCP with higher oestrogen dose</td>
</tr>
<tr>
<td>Saquinavir (SQV)</td>
<td>Oestradiol levels may be decreased; SQV unchanged (unboosted)</td>
<td>Nil documented</td>
<td>As SQV is administered with RTV, caution advised</td>
</tr>
<tr>
<td>Efavirenz (EFV)</td>
<td>Non-significant increase in oestradiol levels; EFV unchanged</td>
<td>Not studied</td>
<td>COCP can be used but advise additional method until interaction fully elucidated</td>
</tr>
<tr>
<td>Nevirapine (NVP)</td>
<td>Significantly reduced oestradiol levels</td>
<td>Significantly reduced norethisterone levels</td>
<td>COCP not advised for contraceptive purposes; increase dose if for other indications</td>
</tr>
</tbody>
</table>


*APV is usually administered as a prodrug now (fosamprenavir) but there are no data currently available on the interaction, if any, between fosamprenavir and hormonal agents.

**Much higher RTV dose than that used for boosting (usually 100 mg once or twice daily, occasionally 200 mg twice daily). COCP, combined oral contraceptive pill.

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Advocating a reduction in injection interval from 12 to 10 weeks in women on PI- or NNRTI-containing HAART is common practice but there is no evidence that this practice is necessary, even when using a potent enzyme-inducer such as rifampicin. While these data suggest that this may not be necessary with NNRTIs, only nelfinavir was studied, an unboosted PI. Current recommendations are to use PIs boosted with low-dose ritonavir and until more data are available a cautious approach is advisable.

Contraceptive implants

Subdermal implants provide a steady release of low-dose progestogen. There is now only one type available, the etonogestrel-releasing Implanon, which provides effective contraception for 3 years. The levonorgestrel-releasing Norplant has been discontinued.

Efficacy is high; combined data from more than 1700 women using Implanon revealed a Pearl index of 0.0 (95% CI 0.0–0.0) with no pregnancies during over 4000 woman-years of use. Implants, unlike injectables, have the advantage of rapid return to normal fertility after the rod has been removed.

However, unlike the depot injectables, current Faculty of Family Planning and Reproductive Health Care (FFPRHC) guidance is that an additional method of contraception should be combined with a subdermal implant during, and 4 weeks after, the use of enzyme-inducing drugs.

A study in Thailand evaluated Norplant in 88 HIV-infected women in the postpartum period and at 24 weeks’ follow-up the method was reported to be safe and well-tolerated. In addition, subdermal implants have been shown to be comparable to tubal ligation in terms of patient satisfaction amongst women wanting a long-term method of contraception.

Emergency contraception

Levonelle is a progestogen-based emergency contraceptive (two tablets of 750 μg levonorgestrel) that can be prescribed or purchased over the counter. WHO data support the use of two tablets immediately as opposed to the traditional advice of taking one tablet followed by the second 12 hours later.

There is evidence that many HIV-infected women are unaware of the availability of Levonelle, and all those who rely on barrier methods alone should be counselled about the availability and window of efficacy of this method.

Current FFPRHC advice is that women on enzyme-inducing drugs should take two tablets as soon as possible followed by a third tablet after a 12-hour interval, although this is outside the product licence and no evidence based. It would therefore be prudent to extend this advice to women on HAART unless evidence to the contrary transpires.

An alternative method of emergency contraception is a copper intrauterine device (IUD), which can be inserted up to 5 days after the first episode of unprotected intercourse or up to 5 days after the earliest predicted date of ovulation. The FFPRHC does not advise routine administration of antibiotic prophylaxis at the time of emergency IUD insertion; however, as HIV-infected women are only WHO Category 2 for IUDs after a negative STI screen, prophylaxis is sensible in this group.

Intrauterine devices

Despite theoretical concerns, prospective data from a Kenyan study showed no difference in infection rates between HIV-positive and HIV-negative women. However, the FFPRHC advises that as long as risk assessment and
screening for bacterial STIs is carried out, HIV-positive women in the UK can be fitted with an IUD. In practice, a low threshold for the use of prophylactic antibiotics is advised. Again, barrier contraception is advised to prevent HIV transmission between discordant couples or of resistant strains between sero-discordant couples.

**Copper IUDs**

These provide a highly effective method of contraception with a Pearl index of 0.6–0.8 in the first year.

**Levonorgestrel-releasing intrauterine system**

The intrauterine system (IUS) is basically a plastic IUD containing a reservoir of levonorgestrel. There is no evidence of a reduction in efficacy of this method with the concurrent use of enzyme-inducing drugs. This study was carried out in 56 women predominantly on anti-epileptics; anti-retrovirals have yet to be studied. The intrauterine levels of progesterone achieved with an IUS are about 1000 times greater than those seen with depot progestogen methods, therefore an IUS is highly likely to remain efficacious in women on HAART.

**Sterilisation**

Sterilisation is the commonest method of contraception for the 40+ years age group with approximately equal proportions of males and females undergoing the procedure. Although highly effective, failure can occur; the lifetime risk of pregnancy following tubal occlusion is 1 in 200 and after vasectomy is 1 in 2000.

Sterilisation should be viewed as a permanent procedure; however, reversal of tubal occlusion and vasectomy is possible. Women should be counselled about the risk of laparoscopic procedures and the possibility of progression to laparotomy. Vasectomy complications are infrequent and long-term sequelae such as chronic testicular pain are rare.

Importantly, vasectomy does not reduce the risk of viral transmission from an HIV-infected male, HIV levels remain the same before and after vasectomy.

**What needs to be done?**

There is a paucity of data on the interactions between anti-retrovirals and hormonal contraceptives. Perhaps more importantly, there are even fewer data on whether any of the actual, or theoretical, interactions translate to a reduction in efficacy in practice. There is therefore an urgent need for more pharmacokinetic trials and long-term follow-up of HIV-infected women using hormonal methods.

Both HIV infection itself and anti-retrovirals contribute to an increased risk of cardiovascular disease. Further data are needed to confirm whether this should be accounted for when considering combined hormonal methods in this population.

In addition, there is a theoretical disadvantage in using high-dose progesterogen-only methods in a group of women already at increased risk of osteopenia and osteoporosis. Once again, further studies are warranted.

**Conclusions**

In the era of HAART, HIV-infected women should be assumed to have the same fertility as their HIV-negative counterparts, both in terms of ability and desire to conceive. There are two important HIV-specific factors to consider: first the need to prevent HIV transmission and second the risk of drug–drug interactions between hormonal contraceptives and HAART/other agents.

Ideally a reliable barrier contraceptive should be combined with a hormonal method to ensure high protection against pregnancy should the barrier method fail.

HIV-infected women need not be denied IUD/IUS methods if screening for, and treatment of, bacterial STIs takes place.

Individuals on HAART should be counselled as to the possibility of detectable HIV in genital secretions despite an undetectable plasma viral load. Sero-discordant couples wishing not to use barrier methods should only do so in consultation with their HIV physician.

Finally, every possible measure to increase the uptake of HIV testing and the rates of disclosure to sexual partners should be undertaken. Only by reducing stigma and the pool of both undiagnosed and undisclosed infection can the risks of transmission be truly minimised.

**Statements on funding and competing interests**

Fundings. None identified.

Competing interests. None identified.

**References**


The David Bromham Annual Memorial Award

David Bromham was the first Chairman of the Faculty of Family Planning and Reproductive Health Care. He died in office in 1996. Throughout his life, David was an energetic and inspirational man. Whilst in Leeds, he set up an assisted conception programme, which was and is one of the most successful in the world. In 1991 he set up a fertility control unit designed to provide a more accessible service for the termination of pregnancy. He also carried out an extensive programme of research and was closely involved with the British Journal of Family Planning (now the Journal of Family Planning and Reproductive Health Care).

The Award is not intended to be a prize for long and distinguished service, rather for a piece of work which through inspiration, innovation or energy has furthered the practice of family planning and reproductive health care in any way and any setting. It is not a recognition grant. Younger health professionals sometimes undervalue their achievements but they are exactly the people that David Bromham would have wished to see encouraged as this award now acknowledges.

The award will be made either to an individual (who must be a current Diplomate or Member of the Faculty) or to a team, which could be multidisciplinary. In the latter case, the lead doctor should be a current member of the Faculty. You may nominate yourself or your team or be nominated by someone else. The award itself, which will be presented at each year’s AGM, will comprise a monetary sum and inscribed memento.

Nomination is by completion of a form that can be downloaded from the Faculty website at www.ffphc.org.uk. Completed submissions must be received at the Faculty office by 10 April annually.

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The Faculty Officers will consider applications for the award and make a recommendation to Faculty Council. Applications for the scholarship are restricted to members of the Faculty.

Applications should include the following details: country and establishment(s) to be visited, aims of visit, details of visit and benefits, together with a brief curriculum vitae and full contact details.

Enquiries about the Scholarship and submissions should be e-mailed to: fulden@ffphc.org.uk.

Completed applications must be received at the Faculty office by 1 April annually.

Entries should be submitted to: International Travelling Scholarship, Faculty of Family Planning and Reproductive Health Care of the RCOG, 27 Sussex Place, Regent’s Park, London NW1 4RG, UK.