HIV transmission in serodiscordant heterosexual couples

Risk is not zero but is low if the infected partner takes antiretrovirals

Antiretroviral treatment inhibits HIV viral replication and reduces plasma viral load. Low plasma viral load is associated with a lower probability of HIV transmission, which opens the possibility of treatment to reduce HIV transmission.1, 2 The feasibility, potential effectiveness, and risks of such treatment are unclear, and a key uncertainty is the extent to which successful antiretroviral treatment reduces HIV infectivity. In the linked observational study, Del Romero and colleagues estimate the risk of heterosexual transmission of HIV-1 from infected people taking combined antiretroviral treatment.3

Only randomised controlled trials comparing transmission in HIV serodiscordant couples, where the infected (index) partner receives or does not receive antiretroviral drugs, can accurately estimate the effect of such treatment on infectivity. The HPTN-052 trial is the only ongoing trial of this type. In this trial, partners infected with HIV are assigned to immediate antiretroviral treatment or deferred treatment when their CD4 count drops below 250 cells/μl. This trial will hopefully provide a good estimate of the effect of antiretroviral treatment in patients with more than 250 CD4 cells/μl, but it will obviously not obtain an estimate in patients with fewer than 250 CD4 cells/μl.

Evidence that antiretroviral treatment reduces sexual transmission of HIV comes from observational studies of serodiscordant couples and ecological studies,4, 5 but transmission from patients on antiretroviral treatment has also been documented.6 Del Romero and colleagues’ study is a welcome addition to the observational studies that estimate infectivity in serodiscordant couples. In partners of patients with HIV not receiving treatment, the incidence of HIV was 0.6 per 100 couple years, whereas none of the partners of patients on antiretroviral treatment seroconverted. The authors also computed the probability of transmission per unprotected sex act (figure). The infectivity estimates were low and imprecise, with no significant difference between treated and untreated patients (figs 1 and 2, see bmj.com). The authors therefore conclude that transmission of HIV from successfully treated patients cannot be excluded, because it is biologically possible and because the data are consistent with one infection per 91 couple years (compared with one per 71 couple years in untreated couples). Moreover, because this is an observational study, the treated and non-treated groups are most probably not comparable. Patients receiving treatment are necessarily different because of their poorer clinical history. Some differences were also seen in risk behaviour and characteristics of the index partners at baseline and during follow-up, which may have increased transmission in couples with an untreated index partner.

How do the results of Del Romero’s study compare with other studies? Apart from four studies from high income countries conducted before the era of antiretroviral treatment that also reported no seroconversion, the infectivity estimates in the non-treated group were generally lower in Del Romero’s study than in previous studies.7, 8 A recent systematic review of observational studies in heterosexual serodiscordant couples found five studies reporting HIV infectivity estimates according to treatment. In two of these studies with information on viral load, no seroconversions were seen when treated index patients had viral loads of fewer than 400 copies/ml. One study from Uganda reported no seroconversions in partners of treated cases despite only 79% having achieved a viral load fewer than 400 copies/ml six months after the start of treatment. A more recent study (not in the systematic review7) in seven African countries observed one seroconversion in 256 person years of follow-up in partners of index cases.

Together, these seven studies of patients receiving antiretroviral treatment independently of viral load will help to provide a more precise overall estimate of the seroconversion rate.9, 10 Although the probability of transmission during antiretroviral treatment seems to be greater than zero, these studies support the idea that treatment reduces infectivity, which could translate into benefits at the population level, as long as risk behaviour does not increase. To date, some studies,3 but not all,10 suggest a beneficial effect of antiretroviral treatment at the community level.

What effect do these data have on counselling serodiscordant couples? In 2008, the Swiss Federal AIDS Commission released a controversial statement to the effect that HIV infected people with undetectable concentrations of the virus (<40 copies/ml) for at least six months, who adhered to a strict antiretroviral drug programme and had no other sexually transmitted infections, were not infectious to their regular heterosexual partners.11 The challenge for counsellors is to ensure that people understand the exact set of conditions and time period when it seems to be safe to have unprotected sex. Del

Transmission risk per sex act

Per sex act heterosexual transmission probability estimates with and without antiretroviral treatment (ART)2 compared with pooled estimates before the ART era.1 n=number of studies included in pooled estimates; FM=female to male, MF=male to female; C=FM and FM combined

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Wellbeing in the workplace

Lack of precise measurement or recommendations should not deter employers from taking action

In November 2009, the National Institute for Health and Clinical Excellence (NICE) published guidance for employers on promoting mental wellbeing through productive and healthy working conditions. Excess work related stress harms employees’ physical and mental health. From an economic perspective, impaired efficiency at work associated with mental health problems costs the United Kingdom £15.1bn (£16.9bn; $22.2bn) a year. From a health perspective, stress at work is consistently associated with increased total mortality and acute myocardial infarction.

Ideally, the guidelines should outline discrete steps that could easily be implemented, improved the efficiency and satisfaction of workers, and ultimately be shown in partnerships with them. This approach should integrate mental wellbeing into all policies and practices concerned with managing people, including those related to employment rights and working conditions.

Unfortunately, some of the advice is so general that it is almost useless. For example, the first recommendation in the guidance is to: “Adopt an organisation-wide approach to promoting the mental wellbeing of all employees, working in partnership with them. This approach should integrate the promotion of mental wellbeing into all policies and practices concerned with managing people, including those related to employment rights and working conditions.” This reads more like a mission statement than a discrete step that a responsible business manager could implement. Other recommendations are more practical, however; examples of how to monitor mental wellbeing (such as attitude or satisfaction surveys, and data on absence rates and employee turnover) are reasonable and appropriate for different sized businesses. Other advice, such as allowing workers flexible hours, and that managers “respond with sensitivity to employees’ emotional concerns” may seem to lag behind policies already in place.

Perhaps the lack of concrete guidance comes from the diverse and changing nature of stress. Stress is not easy to measure—after all, it is subjective. The NICE guidance defines stress as “the adverse reaction people have to excessive pressure or other types of demand placed on them,” meaning that, by definition, workplace stress is excessive. Other major causes of chronic disease—physical inactivity, other recommendations are more practical, however; examples of how to monitor mental wellbeing

Romero and colleagues are more cautious and continue to promote the use of condoms regardless of viral load; in their study condom use reduced HIV transmission by 93%.

On the basis of current evidence, we conclude that although taking antiretroviral treatment reduces the risks of vaginal intercourse with an HIV positive partner, intercourse is not totally risk free. Despite their limitations, additional studies in discordant couples are needed to estimate the infection risk more precisely, especially for homosexual and heterosexual anal intercourse. It is unclear how these results for heterosexual populations will translate to homosexuals.