Evaluation of behavioural interventions in HIV/STI prevention

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Introduction

Behavioural interventions in HIV/STI prevention are a subject of considerable interest and debate. In the absence of effective vaccines such interventions are key to HIV/STI control strategies, but the question of what actually works remains a challenging one. Published research aiming to evaluate the effectiveness of behavioural interventions has been reviewed and mostly found wanting. This article considers why the evaluation of behavioural interventions is often complex, and attempts to unravel some of the surrounding controversy. Questions include what is meant by behavioural interventions? how can they be evaluated? and what can they be expected to achieve in HIV/STI prevention?

What is meant by behavioural intervention?

Various terms, including behavioural, psychosocial, and lifestyle, are used to describe very different kinds of interventions designed to change a wide range of human behaviours. For example, the intervention may range from a brief exchange of information or advice to long term, intensive psychological “counselling” or therapy. And the aim may be to change smoking, exercise, dietary, or sexual behaviour of individuals, small “high risk” groups, or whole communities. In the context of HIV/STI prevention, a behavioural intervention is one that seeks to reduce the risk of acquiring or passing on HIV or other STI by changing behaviours that lead to transmission of infection, principally sexual and injecting behaviour. This still encompasses a wide range of possibilities, as the link between behaviour, change and transmission of infection may be fairly direct (for example, consistent use of condoms between known HIV discordant sexual partners) or much more indirect (for example, raising self esteem or negotiation skills among sexually inexperienced young people to reduce the likelihood of high risk sexual behaviour in the future).

Behavioural interventions have been based on a number of psychological models such as theories of reasoned action, self efficacy, and readiness to change. For example, the model of Prochaska and Diclemente asserts that a series of discrete changes to thoughts and actions are required before behaviour change can be achieved. Whatever the underlying psychological model, the development of a promising behavioural intervention requires careful exploratory research into the determinants of sexual behaviours and the cultural context, values, beliefs, and community norms of target groups. This is the role of qualitative and psychological research which is crucial to the design and implementation of promising interventions but beyond the scope of this brief article. The following focuses on how to evaluate the impact of behavioural interventions in HIV/STI prevention, with particular reference to the role of randomised trials.

How can behavioural interventions be evaluated?

Considerable debate still centres on whether the randomised controlled trial should be considered the gold standard for evaluating the impact of behavioural interventions. It has been claimed that “traditional experimental methods are often hopelessly inapplicable to studies of risk behaviour . . . and of limited feasibility in the evaluation of fledgling community public health programmes.” By contrast, the evidence based approach to the evaluation of healthcare interventions views the randomised trial as the optimal research design for this purpose because of its ability to minimise bias and avoid false conclusions about what works and what does not. To accept the randomised trial as the gold standard, however, is not to deny its limitations and challenges, particularly in the behavioural or psychosocial field.

A frequent objection to randomised trials of behavioural interventions relates to the ethics of “withholding” the intervention from a control or comparison group. This is reminiscent of the debate that flourished when new therapies were increasingly subject to clinical trials in mainstream medicine. Now it is well accepted that anti-HIV therapies should be evaluated in clinical trials before being made available to all who might benefit. Whether it is a drug or a behavioural intervention, the ethical argument for randomised trials rests on “equipoise”—that is, genuine uncertainty that the intervention will actually result in more good than harm. A good example is school sex education, where there are few rigorous studies of the long term effects on the sexual health of young people, but entrenched views among professionals and the general public that school sex education has a positive, negative, or simply no effect on young people. Without well designed trials of sex education, uncertainty and dogma will frustrate efforts to seek a better way forward. Faced with the enormity of the HIV epidemic and the urgent need to find effective behavioural interventions to combat its spread, there has been a tendency to think that action (implementing behavioural interventions) must be preferable to inaction, but
this is essentially a leap of faith rather than a scientific approach, and experience shows that well meaning measures may not work as intended.11

It has also been argued that randomised trials are not appropriate for evaluating behavioural interventions because they ignore the complexity of behavioural and psychosocial interventions.9 Behavioural trials clearly differ in this respect from clinical trials in which the intervention is a new drug whose dose and route of administration have already been established in phase I and II drug trials. By comparison, behavioural or psychosocial interventions are more likely to resemble a “black box”, in which the “active ingredient” has not be identified at the outset. For example, if a support group based on cognitive behavioural therapy proves effective in reducing sexual risk behaviour, is it the cognitive behavioural component of the treatment that works, or is non-specific group support more important? To distinguish confidently between these two possibilities, each component would need to be tested in additional experiments, or in a single trial with multiple arms. Either option is often too costly or lengthy to be realistic. A decision must therefore be made at the outset whether or not to conduct a pragmatic trial, in which the whole intervention package is tested to see whether it advances current healthcare provision, regardless of identifying the active ingredient. Despite some clear differences between clinical and behavioural intervention trials, the ethical principle of establishing that an intervention does more harm than good is unaltered by its complexity. There are plenty of examples of complex chemotherapeutic regimens, for example, that are routinely subject to randomised trials. Perhaps the greatest challenge, in the behavioural field, is to design and conduct trials in such a way that the delivery of the intervention can be standardised as much as possible and the black box made more transparent, so that the benefits of behavioural interventions, once demonstrated in a controlled trial, can be replicated in real life.

**What can behavioural interventions achieve in HIV/STI prevention?**

A key issue here is the choice of outcomes by which to assess the success of a behavioural intervention.12 The ultimate goal of such interventions is to reduce the rate of new HIV/STI infections in defined groups or populations, and many innovative interventions have been set up throughout the world with this goal in mind, but no randomised trial has yet reported the impact of a behavioural intervention on HIV incidence.13 This partly reflects the fact that HIV incidence is relatively low, even in high risk groups (for example, around 5% per year in HIV discordant heterosexual couples,14 or 4% in homosexual men attending central London genitourinary medicine clinics15). Large, costly trials are needed to reliably detect an epidemiologically or clinically meaningful difference (between intervention and control groups) in an uncommon endpoint like incident HIV infection.

Acute (non-HIV) STIs are suitable endpoints for behavioural intervention trials because they are important causes of morbidity in themselves, they occur more commonly than HIV infection, and have been clearly shown to increase the risk of HIV transmission.16 17 In the context of a controlled trial, care needs to be taken that the opportunity for diagnosis of STI is the same in intervention and control groups, otherwise any difference at the end of the trial may simply reflect greater efforts to diagnose STIs in one group than another.

Should studies rely on self reports of high risk sexual behaviour—for example, unprotected intercourse, as outcomes by which to assess the impact of behavioural interventions? Much has been written about the validity of self reported sexual behaviour which cannot be observed or verified directly, and the potential for social desirability bias.18 In a controlled evaluation trial, it is important to be aware that the way in which informed consent is obtained can increase the potential for such bias, particularly where “blinding” to the intervention of interest is not possible. This emphasises the importance of conveying equipoise (see above) to potential participants at the outset, so as to minimise inaccurate reporting of the desired outcome equally across comparison groups. The problem of social desirability bias can also be addressed by new data collection methods. Recent studies in men and women comparing computer assisted self interview (CASI) with conventional pen and paper interviewer questionnaires (PAPI) have shown that disclosure of sensitive or stigmatised behaviours, such as sex with prostitutes, is more likely to occur with CASI.19 Despite these methodological advances in improving the quality of self reported behavioural data, it seems wise to include objective outcomes of high risk sexual behaviour, such as STI markers, wherever possible.

A criticism frequently levelled at behavioural intervention trials is that they do not reflect the situation in real life.13 20 Rarely, however, do interventions of any sort have such a dramatic impact that their usefulness can be established outside a controlled research environment. Despite this, pragmatic trials can be designed to mimic real life conditions as closely as possible without compromising the scientific need to establish a link between intervention and specified outcome(s). Another issue closely related to “real life relevance” is the generalisability of behavioural interventions. Much is rightly made of the need to develop interventions that are culturally sensitive and appropriate to defined target groups. Features of a promising intervention for commercial sex workers in Bombay will obviously differ from those of an intervention aimed at homosexual men in London. But if an intervention is shown to be effective in a particular group, such as homosexual men in clubs and bars in the mid western United States,21 or school students in Uganda, how far can the findings be generalised to others? The answer depends on judgment rather than statistics, so the question of generalisability should be considered early
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