HIV TESTING AND ARV PROPHYLAXIS FOR NEWBORNS WITHOUT THEIR MOTHERS’ CONSENT

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Criminal law, constitutional rights and medical ethics (not forgetting common sense) can at times contradict each other, putting medical professionals on the spot. This article is based on a case study discussed on the HIV Policy & Ethics Discussion Forum: http://groups.google.com/group/policy-ethics.

CASE STUDY

A paediatrician is called to the nursery ward of a government hospital to see a male infant born 8 hours previously. The infant’s mother is 33 years old, wasted and has oral thrush. This is her second child, the first having died in infancy after a short illness with a history typical of pneumonia.

The mother was not offered an HIV test during pregnancy as the clinic she attended did not have such services. A nurse calls the paediatrician as her offer of HIV testing to the mother has been declined. She requests the paediatrician to convince the woman to test, given the benefits that such knowledge gives the woman, as well as to enable the provision of post-exposure prophylaxis for the newborn and of infant feeding counselling. The paediatrician examines the newborn, who is vigorous, fully grown for age and has no signs of HIV infection. She then carefully counsels the patient, explaining the potential harm of testing, and the benefits of HIV testing, for the woman and her infant. The woman still declines.

The paediatrician is aware of the efficacy of antiretroviral (ARV) prophylaxis given to HIV-exposed newborns whose mothers did not receive ARVs.1-3 The former’s conscience and medical duty to act in the best interests of her patient (the child) have to be balanced against hospital and international policies which state that newborns cannot be tested for HIV exposure and be given prophylaxis without their mothers’ consent. She thinks of many other colleagues – such as the previous medical superintendent of the East London Hospital Complex4 – who in similar situations acted from their conscience, even if such actions were contrary to prevailing policies and protocol. The paediatrician then tests the infant, whose antibody rapid tests show he is HIV-exposed. The doctor provides ARV prophylaxis to the infant, counsels the woman about her own HIV status and enrols her in an HIV clinic which provides antiretroviral treatment (ART).

Questions for discussion
1. Was the paediatrician correct to test the infant without the mother’s consent? What is the optimal balance between a woman’s right to autonomy and choice, and her infant’s access to health care services?
2. Was the paediatrician correct to provide ARV prophylaxis to the infant without consulting the mother? Should the paediatrician have informed her that she had given the infant ARV prophylaxis?

DISCUSSION

AN ETHICAL AND RIGHTS-BASED APPROACH

A woman’s constitutional rights to privacy, reproductive choice and bodily autonomy are all too often violated and require adequate legal protection. Also, a woman’s right (and legal obligation) to make choices for her child is common practice. However, HIV infection in infants and its concomitant cost and suffering are essentially preventable. In such circumstances, rights compete and need to be carefully weighed. Dedicated efforts, which are culturally appropriate and, ideally, communicated in patients’ home language, are needed to explore and address the underlying reasons why the woman declined HIV testing. In South Africa, ART is becoming increasingly available, and systems are in place to safeguard confidentiality. In such settings, it is difficult to construct a reason for not testing an infant; when the mother refuses, that is more compelling than an HIV-free child.
1. Any invasive medical treatment or test without the study would be essentially threefold: if she were to do so, the legal ramifications for this case would instigate legal action against the pediatrician; but in a comparison, children who have contracted HIV should intervene when these are undermined. Another possibility for protecting children’s health and wellbeing, and the African context.

2. The mother would also be in a position to report the paediatrician to the Health Professions Council of South Africa (HPCSA) for unethical conduct.

3. The state must assume ultimate responsibility for protecting children’s health and wellbeing, and should intervene when these are undermined. Another consideration is that children who have contracted HIV could in time argue that, by not having been tested for HIV exposure and receiving ARV post-exposure prophylaxis and a woman’s interests in not knowing her own HIV status. Although legislation, policy, and guidelines emphasise the principle of informed consent, the Constitution trumps these. Section 28 of the Constitution states: ‘A child’s best interests are of paramount importance in every matter concerning the child.’ This clause has been used to assert children’s best interests, as in cases where Jehovah’s Witnesses declined blood transfusions for their children.

Where current practice conflicts with the child’s interests, can health care workers act from their conscience, or is this the sole domain of the courts? Where policy and legislation are outdated and lag behind medical progress, bringing a test case to court could precipitate change. For example, in circumstances where a woman refuses HIV testing after birth, a health care worker could launch an urgent court application to test the infant and provide prophylaxis without the woman’s consent. The authors feel that the matter of a paediatrician launching such a case is long overdue. It could be argued that paediatricians each day make active decisions not to test newborns for HIV exposure, even though testing may be in the best interests of those whom they serve.

Several states in the USA have for almost a decade successfully implemented mandatory testing of newborns and thus provided proof of concept and encouraging safety data; this should justify further investigation in the South African context. The state must assume ultimate responsibility for protecting children’s health and wellbeing, and should intervene when these are undermined. Another consideration is that children who have contracted HIV could in time argue that, by not having been tested for HIV exposure at birth, the health providers who cared for them after childbirth neglected to protect them from HIV infection and did not act in their best interests.

LEGAL IMPLICATIONS

In the above scenario, it is doubtful that the woman would institute legal action against the pediatrician; but if she were to do so, the legal ramifications for this case study would be essentially threefold:

1. Any invasive medical treatment or test without the patient’s consent (in this instance, that of the legal guardian of the infant – her mother) constitutes an assault under South African common law as well as an invasion of personal rights.  

2. It therefore follows that the mother could lay a charge of assault on behalf of her child against the doctor who tested and provided medical treatment to the infant without the mother’s consent.

3. The mother would also be in a position to report the paediatrician to the Health Professions Council of South Africa (HPCSA) for unethical conduct.

However, it is unlikely that such a course of action would succeed in court. In her defence, the paediatrician would be able to argue that the court is under a constitutional obligation to develop common law so as to ‘… promote the spirit, purport and objects of the Bill of Rights’ (section 39(2) of the Constitution) and in line with the paramount place given to the interests of the child (section 28(2) of the Constitution).

Evidence is overwhelming that it is not in the best interests of a child to acquire HIV from the mother, and that providing HIV testing and post-exposure prophylaxis will reduce the risk of the child contracting a chronic and life-threatening illness. Moreover, medical evidence shows that administering a single dose of ARVs to an infant is not harmful.

The paediatrician could therefore argue that courts are constitutionally obliged to develop the common law of assault to exclude instances of beneficent intervention in the interests of a minor. With this approach, it is likely that the doctor would be acquitted of a charge of assault, while the Health Professions Council would probably make a similar finding.

CONCLUSION

A test case may effect policy change, though must never negate or minimise the real difficulties that women face in this epidemic, and their needs for care and support. Ideally, women should be strongly encouraged to test and be referred to appropriate programmes during or prior to pregnancy. In lieu of this, the infant’s interests in not contracting HIV are paramount.

Perhaps the epidemic could be reversed with more vigorous interventions, carefully considered and with specific efforts to minimise any human rights infringements. Where access to HIV treatment and confidentiality are assured, the degree and range of benefits of an early HIV diagnosis differ markedly from those of a late diagnosis when HIV inevitably declares itself with severe diseases. Perhaps health workers have for too long protected people from facing an inevitable diagnosis, rather than protecting adults’ health and that of their children. We can never turn back the clock, but we can alter the speed of its ticking.
As it stands, this argument for mandatory newborn testing unnecessarily pits the interest of mother and child against each other and creates conflict where there should be collaboration. Our main concern is that it fails to acknowledge the consequences of the fact that newborn testing amounts to ‘proxy testing’ of the mother – in this case, without her consent. Not only do the paediatrician’s actions violate the mother’s right (and indeed legal obligation) to make medical decisions for her minor infant, they also violate her constitutional rights to privacy, reproductive choice and bodily and psychological integrity. The policy change that Chersich and Richter are urging would disempower and undermine women’s agency on a number of levels. Further, proxy testing of the mother in this way is a violation of fundamental rights that are now recognised and widely accepted as necessary components of ethical HIV diagnosis and treatment. These include the right to informed consent and not to be tested against one’s will. Such rights are enshrined in both national and international policy, guidelines and legislation13-15 and – perhaps most importantly – the South African HIV & AIDS National Strategic Plan 2007 – 2011.16 They are also recognised internationally as good public health practice by WHO and UNAIDS.15,16

The authors admit that mandatory testing could prompt an ‘eroding [of] the right to refuse testing in [other] situ-

15. Stoffberg & Elliott 1933 CPD 148.

**REFERENCES**


**REBUTTAL**

**COERCIVE POLICIES DO NOT MAKE FOR BETTER HEALTH OUTCOMES**

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Although we agree that more culturally sensitive efforts are needed to better understand the reasons why women in such situations may decline testing, much of this is already known. There has been extensive social science research on stigma, denial and blame in the epidemic – and on the role that gender plays in the particular configurations of these collective responses. We also know from studies of HIV disclosure, for example, that the diagnosis itself is still received by many with profound dread. Suicide ideation following a positive diagnosis is common. But it remains the case that women, in particular, bear the brunt of this stigma: they are often blamed for bringing HIV into the home or into a relationship, they face the very real danger of being beaten by an abusive partner, abandoned, shunned, ejected from the home and rendered destitute. These are not uncommon consequences of the abjection that HIV continues to signal for many people, and which makes an HIV diagnosis something to fear and avoid, both for the individual concerned and for the wider community in which they live. Notwithstanding the limited gains made in reducing stigma in recent years, we should not mandate proxy HIV testing for women unless these issues have been more fully addressed.

The woman in the case study cited above may be faced with further challenges. Her own health seems precarious and after giving birth, she might not have had the emotional or physical resources to cope with a positive HIV diagnosis, much less to deal with the implications of her child receiving antiretroviral prophylaxis. Under such circumstances, what chance does the health of the infant have? With no acknowledgement of the known relationship of infant survival to its mother’s wellbeing and survival, the debate is reduced to a simplistic contest between mother and baby. Yet a newborn does not exist in a vacuum; the mother’s health and wellbeing are central to efforts to improve infant health. Furthermore, the paediatrician may have conflicting moral obligations
between baby and mother, but the hospital itself has an obligation to both the mother and the baby. One cannot be ignored at the expense of the other.

This raises a further concern for us. The case study notes that ‘the mother was not offered an HIV test during pregnancy as the clinic she attended did not have such services’. This suggests that the problems need to be addressed upstream, with a particular focus on prongs 1 and 2 of the World Health Organization prevention of mother-to-child transmission (PMTCT) strategy. Thus the first points of intervention would be: helping women in high prevalence regions to assess their own risk of infection, empowering them with knowledge to protect themselves, preventing unwanted pregnancy in women with HIV, and making safe abortion readily available. Then we would need to ensure that all antenatal clinics do have VCT services – and that the quality of counselling and follow-up support is high, so that more women choose to be tested during pregnancy and enter PMTCT programmes if necessary. Community-based interventions, in particular ‘mother-to-mother’ support groups or one-on-one counselling, are powerful and effective.19,20 We suspect that the woman in the case study might have responded differently if she had been counselled by a peer who shared her language and cultural background, and who had perhaps been through similar experiences herself. This would have been preferable to being ‘convinced’ to test in a time of stress where informed consent could not be assured and thus was ignored by a paediatrician whose main concern was clearly the health of the infant. Moreover, consent is important not only from a human rights perspective but also from a medical point of view: when people’s choices are disregarded and when their buy-in is not secured, treatment and follow-up may be compromised. Bringing a test case to court could potentially undermine precisely the purpose it is meant to serve, namely protecting the health of infants. It could also have longer-term public health consequences, since this kind of legal action could deter vulnerable women from seeking out antenatal care at all. The policy and legislative changes proposed by the authors, we argue, are premature if not completely unnecessary.

The authors posit that ‘[i]t is likely to be very small. Indeed, there is evidence that high-quality counselling, uptake of VCT among women in antenatal settings is very high’.22,23 In fact, this vital information is central to the debate – yet the authors make no mention of it. Why, then, the need for a measure as coercive as mandatory newborn testing, enforced by law and policy, when the overall impact of this intervention on the HIV epidemic is likely to be relatively negligible?

Finally, for the authors to invoke the argument about health care workers’ conscience is to assume that decisions made from ‘conscience’ will, in every case, align with what is medically the best decision to make for the patient. But this surely cannot be the case. We have only to consider a comparable situation relating to the implementation of termination of pregnancy policy in South African public health facilities. Here, too, we find health workers acting on the grounds of ‘conscience’ and refusing to have any part in carrying out the procedure. But many of these health workers also refuse to arrange adequate counselling or referral for the women concerned. Acts of ‘conscience’ are admirable, indeed. But they are hardly neutral, objective or necessarily medically correct.

In conclusion, we regard this argument as a classic example of ‘act first, think later’; a narrow, biomedical and legal solution to a complex human problem. We are a long way from ensuring quality services for HIV-positive women and protecting their rights to information, privacy and confidentiality. Our view is that women should not have to pay for the failures of primary HIV prevention and reproductive health services, nor should their rights be sacrificed because political commitment and leadership in the epidemic has been lacking. In short, more debate is needed – and we would urge that such debate involves a wide range of stakeholders: not only maternal and child health specialists and bioethicists, but also experts and advocates in the fields of women’s reproductive health and rights, AIDS activists, civil society organisations, social scientists, and representatives of government. Most importantly of all, we need to hear the voices of ordinary women in South Africa who are actually confronted with such painful dilemmas every day.

REFERENCES


