INTRODUCTION

Sickle cell anaemia (SCA) is the most common form of haemoglobinopathy in Nigeria affecting about 1-3% of the population and it is associated with physical, psychosocial and emotional suffering. Prenatal diagnosis (PND) and genetic counseling are ways of preventing the spread of the disease; however these means of prevention are associated with many ethical dilemmas. Ethical issues discussed in this paper include the safety of the procedures used in obtaining tissue sample for prenatal diagnosis, abortion of affected fetuses and the question of genetic selection. Finally, the ethical implications of genetic counseling and issues relating to the principle of justice in healthcare are highlighted.

ETHICAL ISSUES

The issue of safety of the procedures used for PND is worth mentioning first. Though they are found to be relatively safe, there is still a chance of a miscarriage following chorionic villus sampling and amniocentesis (worse with CVS and usually multi-factorial). Abortion of the affected fetus is regarded as a component of PND in most cases. In the case of the fetus having the SS genotype, the ethical question arises whether to have an abortion or to keep the pregnancy. The decision whether to terminate a pregnancy based on a positive result is usually a difficult one that involves religious, psychosocial and cultural considerations.

Another reason why the abortion of an affected fetus may not be accepted is because of the risk of complications mainly due to lack of reliable and safe health care practices in developing countries. Even when abortion is legal in the local context, the question whether it is right to terminate an innocent life is still a much debated issue. If on the other hand, a decision is made to keep the pregnancy, the question whether it is right or not to bring a child with a disease condition that causes so much suffering to the world. Utilitarians
would argue that it is more cost-effective to abort the affected fetuses as this will reduce on the long run the socio-economic and emotional consequences of the disease. Bringing up the question of cost-effectiveness and life of patients would be looked at in many developing countries like Nigeria as being cold and inhuman, but the reality of scarcity of resources and rationing of health care resources is there for all to see. The introduction of pre-implantation genetic diagnosis (PGD) in developed countries would seem to have put the controversial issue of termination of affected pregnancy at rest as only “genetically healthy” embryos will be transferred to the uterus. The practice itself is laden with its own ethical dilemmas relating to: the moral status and destruction of embryos, tendencies for eugenic practices (selection of particular traits in the embryo), the possibility of long term complications from the procedure and distributive justice issues. Opponents of these prenatal or pre-implantation tests would also argue that SCA is now becoming a chronically manageable disease with increase in life expectancy and quality of life of patients hence there is no reason for these investigations.

Some people would argue that using a prenatal diagnosis for SCA would lead to a systematic elimination of the genetic mutation from the population. Could this be called a form of eugenics? The author definitely does not think so as the choice here is not about specific traits that are desired in a child but having a child free of a particular genetic disorder. The right to know is a fundamental right of the couple hence carrying out a PND for SCA actually empowers the couple to plan for the new child (if they decide to keep the pregnancy – if SS genotype) and gives them peace of mind (if AS or AA genotype). This is the autonomous choice of the couple, a right to decide what is acceptable to them. There is also the risk of pressure being put on the couple directly or indirectly by the society to have prenatal diagnosis done because of the availability of the tests (the so called technological imperative). This could lead to affected couples being blamed for not making use of the tests to avoid having children with sickle cell disease.

The scope of genetic counseling offered by health care workers (clinical geneticist, haematologist, nurse or obstetrician) has a lot of ethical implications. Several studies have clearly shown that genetic counseling is considered as one of the best ways of controlling the disease. As usual with genetic counseling a “non-directive” and “client-centered” form of counseling is preferred. This form of counseling entails disclosure of genetic risk information necessary for the clients to make an informed decision without direct or indirect advice from the counselors. But the question whether it is possible to be absolutely non-directive in counseling is one that is generating a lot of debates among bioethicists and geneticists. How can one ignore his or her own intuition and sense of morality when confronted with challenging situations? Is it possible for a counselor who is a practicing Roman Catholic to be value-neutral when discussing abortion as an option after PND? Counseling for SCA is even more problematic because of the variability in severity of the disease among different patients: while some live fairly crises-free lives for long (50-60yrs), some die during childhood or adolescence. So how do we know which form of the disease the unborn child is going to develop?

Finally, the principle of justice in health care requires that access to PND be fair and equitable. This is definitely not so because of intra-country and inter-country disparity in access to PND; many people requiring prenatal diagnosis for SCA may not have access to it because of lack of the service in their environment or inability to pay for the services. This is particularly common in developing countries of Sub-Saharan Africa where payment for health care is still mainly “out of pocket”. This problem of access to prenatal diagnosis remains one of the major impediments to the control of SCD in developing countries.

CONCLUSION
This review article has clearly shown the ethical dimensions of prenatal diagnosis in the setting of a developing country like Nigeria. It is important for health care providers to be conscious of these issues especially when providing service to clients. There is also a need to build capacity in the area of genetic counseling as a means of controlling the spread of sickle cell anaemia.

REFERENCES


