International Biomedical Research and Research Ethics Training in Developing Countries

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Abstract
Developing an ethical framework for international biomedical research that is receptive to different cultural norms and moral values requires continuous and earnest dialogue between scientists from developed and developing countries. Adequate and sustainable training programs in biomedical research ethics for scientists from developing countries are essential to prepare competent partners from developing countries in this dialogue.

Background
In recent years, international research collaborations have been growing rapidly. Multinational pharmaceutical companies and academic institutions have expanded their research to developing countries. This in part is driven by the increased awareness of health as a global issue, the challenges imposed by new and resurgent global epidemics, the increased demand for new treatments, and practical and financial considerations of the pharmaceutical industry [1,2]. For example, a sharp increase in research in developing countries is largely due to the global HIV/AIDS epidemic and increased efforts to control malaria epidemic [3,4]. In addition, institutions within developing countries are increasingly engaged in biomedical research in order to map local health problems and allocate limited resources to remedy those problems. However, this research is often poorly regulated and does not meet widely accepted ethical standards [5,6].

The globalization of biomedical research – purportedly guided by ethical principles established in developed countries – has led to numerous controversies and ethical dilemmas [7]. For example, the use of placebo controls in clinical trials for preventing perinatal HIV transmission provoked an intense debate concerning issues of research design, exploitation and fairness, and the relationship between a clinical investigator and a human subject [7-9]. This debate involved not only procedural and regulatory questions, such as those related to implementing research protocols or documenting consent, but also deeper questions concerning the social, cultural, and philosophical foundations of research.

Many controversies concerning the ethics of research in developing countries stem from differences in cultural and social norms, health priorities and policies, and wealth disparities between developed and developing countries. For example, informed consent to participate in research depends strongly on the principle of individual choice and autonomy that is deeply entrenched in western culture and philosophy, but is not a central value in many non-western societies [7,10,11]. Consequently, in developed countries, the individual decides whether to participate in research, and that process is documented by signing an informed consent form. However, neither the concept (individual decision-making) nor the procedure (signing a written document) is universally appropriate in traditional societies. In many traditional societies in developing countries some form of family or community consent must be obtained before consent is sought from potential subjects [12,13]. In other words, while community involvement in decision-making may be seen as an infringement of the rights of affluent, well informed individuals living in developed countries, it can be viewed as a layer of protection by poor and illiterate individuals living in developing countries [14,15]. This culturally-based dissociation between group-centered decision-making in traditional communities and the emphasis on the autonomous individual in Western guidelines of informed consent process has greatly complicated the implementation of the informed consent principle in the realities of international research [16,17]. Researchers quickly discovered the urgent need to reconstruct the standards of informed consent in a flexible framework that is more accommodating to cultural diversities [16,18].

Likewise, these difficulties in applying Western guidelines to biomedical research in developing countries are also relevant to other aspects biomedical ethics, including the principles of beneficence and justice [19].

Ethical Relativism, Ethical Universalism and Ethical Pluralism

Anthropological studies suggest that there are deep and widespread moral disagreements across different societies [20]. Practices and actions that are regarded as morally unacceptable by one society may be regarded as acceptable by another. Prominent examples include polygamy and arranged marriages. These observations have led some anthropologists and philosophers to conclude that all moral judgments are relative to a particular culture or society [20]. When applied to international biomedical research, this point of view implies that the ethics of a given study cannot be judged only by the standards developed in western cultures. This is because it is not possible to objectively determine the legitimacy of one culture’s value system over that of another [21]. One consequence of this position is that ethical systems of host communities should have equal force to Western guidelines in governing the conduct of international research [21]. For example, if a non-western society requires that women participate in research only if they have spousal consent, but also deeper questions concerning the social, cultural, and philosophical foundations of research.

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approval, then westerners conducting research in that society should honor this practice. In fact, some commentators and political leaders have argued that insisting that research conducted in developing countries meet the ethical standards of developed countries is a form of moral imperialism [7,22]. Some authors even argue that insisting on seeking individual informed consent to participate in research in certain communities in developing countries, may weaken the social fabric of the society and introduce disorder in it with long-term consequences [23].

On the opposite side of the debate, some ethicists maintain that ethical standards are universal and must be applied no matter where the research is being performed [24]. Moreover, they argue that local laws and traditions cannot justify abandoning those universal principles [25].

Both the relativist and the Universalist point of view are problematic when applied to trans-cultural biomedical research. The former does not provide adequate ethical guidance to conduct research as it does not specify a mechanism to evaluate conflicting cultural views, while the latter does not recognize the huge diversity of cultural norms and ethical expectations in different societies. A middle-ground solution is offered in the form of ethical pluralism, which addresses major problems of the universalist and the relativist positions by acknowledging the central role of local cultures in shaping ethical rules of a given society, and by providing a mechanism to solve ethical disagreements between visiting researchers and host communities through continuous dialogue, mutual evaluation and negotiation [21]. Under the pluralistic approach, negotiated settlements of ethical conflicts provide culturally-relevant guidelines to the specific research project at hand, instead of forcing rigid international ethical guidelines based on unrealistic assumptions of their universal applicability. This is very different from moral relativist approach in that although it recognizes and honors local traditions and practices, it involves a continuous critical appraisal of those values with a vision of a negotiated settlement when ethical conflicts arise. In addition, ethical pluralism contends that culture not only determines the content of the ethical system of a given community, but also its form [26]. That is, local culture determines how ethical values are expressed on the procedural level. This means that in some instances conflicting views may arise on implementation of an ethical principle even though there is no disagreement over the principle itself. For example, in some settings it is in appropriate to ask for signing a printed document after an oral consent has been provided by the potential participant [18]. Here, too, the pluralistic approach envisions negotiated solutions for means to implement ethical standards that are culturally-appropriate.

Therefore, and in order to conduct international biomedical research that is based on universally accepted ethical standards, and takes into account the great diversity in cultural values and social norms globally, we need to develop a trans-cultural ethical framework that is more receptive to variable ethical views of different societies. This framework can serve as general reference for international biomedical research, but which also includes detailed and specific guidelines on how to resolve ethical disagreements based on continuous negotiations and dialogue among the concerned parties: visiting researchers, local scientists and representatives from the host community [27].

**Towards A Trans-cultural Ethical Framework for Biomedical Research**

From this impassioned debate about the ethics of international research, two ideas have emerged that are notable for their relevance and applicability: [1] the need for trans-cultural research ethics, and [2] negotiated or procedural pluralism [21,27,28]. The first notion implies that an ethical framework for biomedical research should take into account the differences in ethical values and social norms across different cultures [29,30]. The second notion builds on the first and recognizes that agreement on universal ethical principles need not constrain the means pursued in following them [31]. That is, different approaches and strategies may equally facilitate the achievement of a single ethical standard or goal [28]. Consequently, useful approaches to research can be reached through negotiations and dialogue between sponsoring agencies and visiting researchers, on one side, and scientists and representative from host communities on the other.

Importantly, the negotiation of ethical guidelines and procedures need not involve compromising established ethical norms in research. Coercion, for example, is never acceptable. Yet what constitutes coercion and what does not, in a particular study and setting, should not be rigidly defined in some office in the country of the sponsoring agency. Rather, it should be a negotiated agreement between sponsors and visiting researchers, and local researchers and host community leaders. Of relevance in this regard is Emanuel et al.’s call to move from principles to benchmarks [32]. They suggest 31 ethical benchmarks which provide a set of practical measures that constitutes an ethical framework for multinational biomedical research (Table 1). These benchmarks can be viewed as blueprints for a negotiation plan to achieve ethical goals in research among the interested parties. A setting-specific version can be developed along the general lines of these benchmarks and negotiated among the sponsors, visiting researchers, local scientists and representatives of the host community.

Fruitful negotiations, however, require trained and competent negotiators on both sides, but little attention is paid to the lack of adequate training and expertise in the area of research ethics of researchers and professionals in regulatory bodies in developing countries [6,33].

**The Need for Adequate and Sustainable Training Programs in Research Ethics for Scientists in Developing Countries**

As international research has evolved, researchers from developed countries have become increasingly dependent on colleagues in the host countries to overcome linguistic and cultural barriers and to help implement research protocols. This assistance does not involve simply a correct translation of study protocols or informed consent documents, but implementing these documents in a culturally and socially appropriate manner. Untrained local researchers, therefore, may jeopardize not only the scientific merit of the study (e.g., by incorrectly translating a questionnaire item), but also its ethical standards. Two possible scenarios might be anticipated in this regard; both are problematic. Because of the strong influence of their own culture and the lack of training in ethical principles of biomedical research, local researchers may unknowingly inject their own values into study protocol and procedures, therefore engaging in a kind of ethical relativism where local norms take precedence over universal ethical principles. Alternatively, local researchers may subscribe entirely to western ethical norms, thus leading to a study that is legally valid, but ethically inappropriate (e.g. asking an illiterate subject to sign an informed consent document that she
Select the study population to ensure scientific validity of the research. Select the study population to minimize the risks of the research. Involve the community in establishing recruitment procedures and incentives. Disclose information in culturally and linguistically appropriate formats. Ensure the freedom to refuse or withdraw. Provide enrolled participants with information that arises in the course of the research.

Scientific validity

Ensure that the scientific design of the research realizes social value for the primary beneficiaries of the research. Ensure that the scientific design realizes the scientific objectives while guaranteeing research participants the health-care interventions to which they are entitled. Ensure that the research study is feasible within the social, political, and cultural context or with sustainable improvements in the local health-care and physical infrastructure.

Table 1: Ethical principles and benchmarks for multinational clinical research.

| Principles                              | Benchmarks                                                                 |
|----------------------------------------|-----------------------------------------------------------------------------|
| Collaborative partnership              | Develop partnerships with researchers, makers of health policies, and the community. Ensure that researchers from developing countries are trained in international ethical principles and regulations. To be able to engage in a mutual evaluation of the local and international ethical standards of studies designed by researchers from other professional in regulatory bodies in developing countries can become an important factor for the sustainability of training programs. |
| Social value                           | Specify the beneficiaries of the research—who. Assess the importance of the health problems being investigated and the prospective value of the research for each of the beneficiaries—what. Enhance the value of the research for each of the beneficiaries through dissemination of knowledge, product development, long-term research collaboration, and/or health system improvements. Prevent supplanting the existing health system infrastructure and services. |
| Scientific validity                    | Ensure that the scientific design of the research realizes social value for the primary beneficiaries of the research. Ensure that the scientific design realizes the scientific objectives while guaranteeing research participants the health-care interventions to which they are entitled. Ensure that the research study is feasible within the social, political, and cultural context or with sustainable improvements in the local health-care and physical infrastructure. |
| Fair selection of study population     | Select the study population to ensure scientific validity of the research. Select the study population to minimize the risks of the research and enhance other principles, especially collaborative partnership and social value. Identify and protect vulnerable populations. |
| Favorable risk-benefit ratio           | Assess the potential risks and benefits of the research to the study population in the context of its health risks. Assess the risk-benefit ratio by comparing the net risks of the research project with the potential benefits derived from collaborative partnership, social value, and respect for study populations. |
| Independent review                     | Ensure public accountability through reviews mandated by laws and regulations. Ensure public accountability through transparency and reviews by other international and nongovernmental bodies, as appropriate. Ensure independence and competence of the reviews. |
| Informed consent                       | Involve the community in establishing recruitment procedures and incentives. Disclose information in culturally and linguistically appropriate formats. Implement supplementary community and familial consent procedures where culturally appropriate. Obtain consent in culturally and linguistically appropriate formats. Ensure the freedom to refuse or withdraw. |
| Respect for recruited participants     | Develop and implement procedures to protect the confidentiality of recruited and enrolled study participants. Ensure that participants know they can withdraw without penalty. Provide enrolled participants with information that arises in the course of the research. Monitor and implement interventions for medical conditions, including research-related injuries, for enrolled participants at least as good as existing local norms. Inform participants and the study community of the results of the research. |

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