Incorporating ethical principles into clinical research protocols: a tool for protocol writers and ethics committees

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ABSTRACT
A novel Protocol Ethics Tool Kit (‘Ethics Tool Kit’) has been developed by a multi-stakeholder group of the Multi-Regional Clinical Trials Center of Brigham and Women’s Hospital and Harvard. The purpose of the Ethics Tool Kit is to facilitate effective recognition, consideration and deliberation of critical ethical issues in clinical trial protocols. The Ethics Tool Kit may be used by investigators and sponsors to develop a dedicated Ethics Section within a protocol to improve the consistency and transparency between clinical trial protocols and research ethics committee reviews. It may also streamline ethics review and may facilitate and expedite the review process by anticipating the concerns of ethics committee reviewers. Specific attention was given to issues arising in multinational settings. With the use of this Tool Kit, researchers have the opportunity to address critical research ethics issues proactively, potentially speeding the time and easing the process to final protocol approval.

The principal goal of clinical research, even when benefiting individual trial participants, is to advance ‘generalisable knowledge’ to help future patients. While that goal is laudatory, clinical research is fraught with ethical challenges including those that occur when research is conducted across multiple trial sites, in different countries or regions, in low-resource settings, in developing countries and with different, sometimes vulnerable, populations. The written clinical trial protocol is the appropriate instrument to illuminate, acknowledge and address ethical challenges specific to each individual study. However, writers of clinical trial protocols—members of the clinical research team in either industry, non-profit or academic settings—may not have access to satisfactory single-source guidance to identify and address relevant ethical issues. The lack of guidance results in clinical trial protocols that either are silent on the ethical issues or include non-specific language in the context of ethical principles without explicitly delineating such principles or challenges. In the absence of explicit description or discussion of ethical issues and choices, ethics committees (RECs), depending on the region, may identify the ethical issues implicit in the clinical trial protocol, infer how protocol writers addressed concerns and may assume—without seeing evidence to the contrary—that ethical issues were not considered and appropriately managed. The lack of explicit description of, approach to and mitigation of ethical issues in a clinical trial protocol can result in time-consuming delay, as ethics committees pose questions that the writers must then answer in a later resubmission. Of even greater importance, not anticipating and planning for important ethical issues may potentially lead to problems in the trial itself.

To provide guidance and to raise the overall quality of clinical trial protocols, the Multi-Regional Clinical Trials Center of Brigham and Women’s Hospital and Harvard (MRCT Center) undertook the initial development of a Protocol Ethics Tool Kit (‘Ethics Tool Kit’), accompanied by a guidance document with points to consider (available in fillable Microsoft Word format at http://mrctcenter.org/resources/2014-11-14-training-material-mrct-ethics-essential-elements-and-points-to-consider-reference-document-toolkit/). The intent of these resources is to help protocol writers recognize and address common ethical challenges in clinical trials, with specific attention to issues that arise in multinational settings. The Ethics Tool Kit is also intended to help ethics committees review and analyse clinical trial protocols in a more efficient, explicit and comprehensive manner.

BACKGROUND
Clinical trial protocols are central to the conduct of clinical trials and facilitate evaluation and review by key stakeholders, including regulators and ethics committees. Despite the importance of sound, well-written and ethical clinical trial protocols, existing

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1The MRCT Center was founded in 2009 to improve the design, conduct and oversight of multi-regional clinical trials, focusing on trials conducted in emerging economies and the developing world. The MRCT Center seeks to establish common, explicit, feasible and ethical standards for conduct of transnational clinical research (http://www.mrctcenter.org).
guidelines for protocol writers have had limitations such as insufficient stakeholder involvement, lack of systematic development and weak empirical support. Two relatively recent documents provide a structure and define needed components of a clinical trial protocol, although neither focused specifically on the ethical issues raised by a planned study. The CONSORT (Consolidated Standards of Reporting Trials) Statement, updated in 2010, presents systematic evidence-based guidance for organising final study reports based on a checklist, and this can be used to inform protocol writing. CONSORT’s checklist highlights 3 of the 11 elements contained in our Ethics Tool Kit, specifically the importance of proper study design, the choice of study population and the criticality of addressing potential harms. Similarly, the SPIRIT (Standard Protocol Items: Recommendations for Interventional Trials) 2013 Statement, generated by an international group of stakeholders, recommends minimum standards for inclusion in clinical trial protocols. SPIRIT includes two topics that might particularly require ethical consideration: the importance of informed consent and trial design, which are both included and broadened in our Ethics Tool Kit. Neither CONSORT nor SPIRIT comprehensively and directly addresses ethical issues in clinical trial protocols. The Ethics Tool Kit complements and expands these two prior documents by offering more focused guidance for identifying and treating ethical issues in clinical trial protocols.

METHODS
The development of the MRCT Center Ethics Tool Kit involved (1) formation of a working group, (2) literature review and (3) review of a sample of 100 approved clinical trial protocols. Following these initial steps, the multi-stakeholder working group aggregated, aligned and reviewed focused ethical questions that were then formatted as (4) an Ethics Tool Kit and accompanying guidance document to allow dynamic usage by protocol writers and ethics committees alike.

FORMATION OF A WORKING GROUP
A group of 20 experts from academic institutions (6), pharmaceutical companies (4), non-profit organisations (4), law firms (3) and ethics committees (3), with backgrounds in clinical trials, medicine, bioethics and law was formed by the MRCT Center in 2012 to create a list of ethical elements that should be addressed when writing and/or reviewing a clinical trial protocol. Each member introduced potential ethical elements by drawing upon the literature and protocol reviews, the MRCT Center work group undertook a descriptive review of a sample of 100 clinical trial protocols to determine if the Essential Elements drafted through consensus were present in current approved clinical trial protocols and, if they were, whether they were discussed directly from an ethical perspective. To minimise bias in the choice of clinical trial protocols to be evaluated and the review process, the review was conducted using a set of predefined guidelines. Protocols were selected based on the following criteria:

- Protocols that had been reviewed and approved by ethics committees
- Multi-site trials with at least one site outside of the USA
- Interventional trials, including medical, social/behavioural and devices
- Trials involving greater than minimal risk, as defined by US research regulations

Selection was retrospective and consecutive from the start date, 30 June 2013, proceeding back in time until 100 clinical trial protocols matching the selection criteria were identified. Prior to selection, no protocol was reviewed for content (other than for the selection criteria listed above). Informed consent forms were also reviewed when available, as some of the Essential Elements might be addressed in the informed consent form instead of the protocol. Two authors reviewed each protocol; if there was disagreement on the assessment, a third author arbitrated. However, little disagreement between the two primary reviewers actually occurred; the kappa statistic, which measures inter-rater agreement, was 0.96.

RESULTS
Recommendations of the working group
Although the assembled working group members had extensive involvement with various aspects of study design and protocol assessment, in their experience, clear and specific discussion of primary ethical issues in clinical trial protocols was unusual. This working group recommended that a dedicated ethics section be included in every protocol. Inclusion of such a section would (1) help clinical research teams proactively consider and articulate ethical considerations associated with their protocol and, as a result (2) improve the dialogue between ethics committees and clinical research teams and among clinical research team members themselves. No working group members were aware of a similar prior suggestion or exposition of how such a section should be structured.

Based on their collective expertise, and informed by the literature and protocol reviews, the MRCT Center work group identified 11 items (called Essential Elements; also see table 1) that should be considered for discussion in a dedicated ethics section within a clinical research protocol.

1. Addressing Relevant Question
2. Choice of Control and Standard of Care
3. Choice of Study Design
4. Choice of Subject Population
5. Potential Benefits and Harms
Table 1  Essential Elements and survey results (sample size=100 protocols)

<table>
<thead>
<tr>
<th>Essential Element</th>
<th>Percent of protocols covering element (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Addressing relevant question</td>
<td>96</td>
</tr>
<tr>
<td>2. Choice of control and standard of care</td>
<td>59</td>
</tr>
<tr>
<td>3. Choice of study design</td>
<td>44</td>
</tr>
<tr>
<td>4. Choice of subject population</td>
<td>39</td>
</tr>
<tr>
<td>5. Potential benefits and harms</td>
<td>76</td>
</tr>
<tr>
<td>6. Informed consent</td>
<td>56</td>
</tr>
<tr>
<td>7. Community engagement</td>
<td>9</td>
</tr>
<tr>
<td>8. Return of research results and incidental findings</td>
<td>49</td>
</tr>
<tr>
<td>9. Post-trial access</td>
<td>22</td>
</tr>
<tr>
<td>10. Payment for participation</td>
<td>40</td>
</tr>
<tr>
<td>11. Study related injury</td>
<td>43</td>
</tr>
</tbody>
</table>

6. Informed Consent
7. Community Engagement
8. Return of Research Results and Incidental Findings
9. Post-Trial Access
10. Payment for Participation
11. Study Related Injury

**Literature review**

Using the search criteria listed above, the literature review found only one relevant scholarly article that provided guidance for drafting and/or reviewing the ethical elements of a clinical trial protocol. The study was published in the psychiatry literature more than 15 years ago, and referenced the Research Protocol Ethics Assessment Tool (RePEAT), a 24-item checklist that contained some of the items identified in our work. Thus, we found little available guidance in a single organised format to guide which items should be considered for discussion in a protocol and how these considerations might be organised in a dedicated ethical section.

**Clinical trial protocol review**

A total of 100 clinical trial protocols were reviewed to determine if the 11 Essential Elements the working group drafted were present in the current approved clinical trial protocols and, if they were, whether they were discussed explicitly from an ethical perspective. A total of 40 clinical trial protocols were identified from publicly available published trials in the *New England Journal of Medicine*, 40 had been approved by independent central IRBs and were available to one of the working group members and 20 had been approved by academic IRBs and were available to one of the working group members. Of the 100 clinical trial protocols reviewed, 37 were funded by industry, 32 by governments, 3 by academic institutions, 3 by different combinations of the above categories and 5 were of indeterminate funding source.

As summarised in table 1, our list of 11 Essential Elements was variably addressed in these 100 clinical trial protocols. For example, while the first element, ‘Addressing Relevant Question’, was almost always included in a clinical trial protocol (96%), other Essential Elements were mentioned much less frequently. It is not surprising that some Essential Elements such as Community Engagement or Post-Trial Access, for example, were mentioned in only 9% and 22% of protocols, respectively. Community Engagement and Post-Trial Access may not be relevant to some protocols, and the latter is, admittedly, an emerging issue. However, evidence of the thinking around Potential Benefits and Harms was not addressed in 24% of protocols, and Challenges in Informed Consent was not found in 44%. Other Essential Elements that might be expected to be important for almost all protocols (Elements 2, 3, 4, 6, 9, 11) were mentioned in 39%–59% of protocols. This variability may not be surprising as no regulation presently requires explicit discussion of ethical issues in written clinical trial protocols or informed consent forms. Absent regulatory requirements, study sponsors and funders may not dedicate resources to document the background thought processes in protocols.

The significance of these findings should not be overinterpreted. The lack of documentation of ethical considerations does not mean that the study was unethical, only that the thinking behind the choices made (eg, in study design, in study population choice, etc.) was not explicit. In addition, it does not imply that the ethics committee did not consider the ethical issues; the ethics committee meeting minutes were not reviewed nor were the exchanges, written or otherwise, between the ethics committee and the principal investigators. Further, the review itself of the 100 sampled protocols had limitations including the small sample size, the admittedly non-representative nature of the protocols that were available and the use of non-validated review criteria. Some clinical trial protocols were analysed by representative(s) of organisations from which the protocols were obtained. This was necessary to protect confidentiality but may have introduced bias into the assessment process. In addition, no regulation presently requires explicit discussion of ethical issues in written clinical trial protocols or informed consent forms. Nonetheless, these findings suggest that critical ethical issues typically of serious concern to ethics committees are often not addressed explicitly in submitted clinical trial protocols.

**The Protocol Ethics Tool Kit: a tool to recognise and address ethical issues**

To ensure and reinforce adequate exposition of ethical issues within clinical trial protocols and to ease the burden of distilling and including this information, the MRCT Center working group developed the Protocol Ethics Tool Kit incorporating the Essential Elements. The Ethics Tool Kit was developed to (1) provide protocol writers and study teams with a tool to recognise and address common clinical trial ethical issues and (2) to

**Table 2  Components of the MRCT Center’s Protocol Ethics Tool Kit**

<table>
<thead>
<tr>
<th>Section</th>
<th>Description</th>
<th>For use by</th>
</tr>
</thead>
<tbody>
<tr>
<td>Short Explanation of the Ethics Essential Elements</td>
<td>A list of the 11 essential elements to consider when writing or reviewing a protocol with an accompanying description of each element</td>
<td>Protocol writers, ethics committees</td>
</tr>
<tr>
<td>Points to Consider</td>
<td>Examples of detailed points to consider for each of the Essential Elements</td>
<td>Protocol writers, ethics committees</td>
</tr>
<tr>
<td>Examples</td>
<td>Examples of language from actual clinical trial protocols that addressed a particular Essential Element</td>
<td>Protocol writers</td>
</tr>
<tr>
<td>References</td>
<td>Relevant citations and sources</td>
<td>Protocol writers</td>
</tr>
</tbody>
</table>

ensure that ethics committees are able to evaluate clinical trial protocols comprehensively and efficiently. The Ethics Tool Kit is not intended to prescribe requirements, to limit ethical considerations or to impose mandates on how ethical issues must be addressed in a trial protocol. Rather, the Ethics Tool Kit is intended to guide thought and discussion and to ensure that ethical concerns specific to a clinical study are, at a minimum, considered in protocol development and made explicit in the protocol itself.

The Ethics Tool Kit is structured in such a way that it can be adapted to meet an individual user’s needs and address specific challenges. Each Essential Element has (1) a short explanation, (2) specific points to consider, (3) background information, (4) practical examples and (5) references. Table 2 provides a brief description of the components of the Ethics Tool Kit and to whom they could be relevant. An online supplementary table S4 presents the short explanation and specific points to consider for each Essential Element. The Ethics Tool Kit in its entirety can be accessed at http://mrctcenter.org/resources/2014-11-14-training-material-mrct-ethics-essential-elements-and-points-to-consider-reference-document-toolkit/.

Use of the Ethics Tool Kit may surface ethical issues that would be otherwise unexplored and also encourage rational, clearly articulated responses. For example, see table 3 on Essential Element 8: Return of Research Results and Management of Incidental Findings. The Ethics Tool Kit is not intended to serve as an exhaustive list of ethical issues that can occur in clinical research, and not every Essential Element is necessarily relevant to every protocol. However, it is recommended that protocol authors consider all Essential Elements, address those that are pertinent for the particular clinical trial and supplement as needed. Authors may choose to discuss ethics throughout the protocol, but the working group sees value in the practice of detailing ethics approaches in a dedicated ‘Ethics Section’ of the protocol.

Computer-based training of the Ethics Tool Kit

In February 2014, in an effort to disseminate the working group’s efforts more widely to researchers in low-income and middle-income countries, the MRCT Center collaborated with colleagues at the Global Health Network at Oxford University, Oxford, England and adapted the Ethics Tool Kit for an innovative digital platform (https://globalhealthtrainingcentre.tghn.org/essential-elements-ethics/). The Essential Elements were first reviewed by our collaborators and then tailored to the electronic format; meaningful visual components were added and assessments at the end of each module were integrated to gauge understanding. A total of 11 course modules corresponding to the 11 Essential Elements have been freely available online through the Global Health Training Centre since 2 December 2014. The uptake of the course has been higher than expected. As of 22 June 2015, the total number of modules taken was 3536 by 1024 users globally. The most commonly accessed modules were (1) Addressing the Relevant Question, (2) Choice of Control and Standard of Care, (4) Choice of Participant Population and (8) Return of Research Results/Incidental Findings. The e-learning course modules were reviewed for content and user-friendliness by the intended end users (investigators in low-income and middle-income countries) and by the leadership of the Global Health Network prior to public release. Furthermore, the Global Health Training Centre e-learning courses are formally recognised for quality and contenti by the Liverpool School of Tropical Medicine (http://www.lstmed.ac.uk/), the Bill and Melinda Gates Foundation (http://www.gatesfoundation.org/) and the Worldwide Antimalarial Resistance Network (http://www.wwarn.org/).

DISCUSSION

The Ethics Tool Kit has potential uses for individual protocol writers and study teams, study sponsors and ethics committees. For individual protocol writers and study teams, it provides a systematic and methodical approach to address the ethical implications of a planned clinical trial. This will assist protocol writers by alerting them to the important ethical issues in study design, enrolment and conduct of clinical trials, and will encourage articulation of appropriate ethical justification. The framework may be particularly valuable to those with less experience drafting clinical trial protocols. The guidance also may be used beyond the protocol, as it can prompt consideration of context-specific difficulties, pertinent policies and local regulatory requirements. For example, the Ethics Tool Kit may alert investigators in low-resource regions to consider challenges in assessing competencies of local sites, differing local medical standards and potential risks of exploitation of local and/or vulnerable populations.

For study sponsors and funders, the Ethics Tool Kit may be useful for documenting the nature of questions that were
considered in protocol design and the analytical approach that formed the basis of the final design. Further, the Ethics Tool Kit may provide sponsors and study teams important insights into the research review process by delineating what research ethics committees are assessing when reviewing studies.

For ethics review committees, the review may be streamlined significantly by altering the protocol model to one in which the ethical reasoning is included in the original submission to the ethics committee. Without an explicit ethics discussion, an ethics committee is left to discern the ethical reasoning behind protocol decisions. When questions arise, the ethics committee engages the principal investigator in dialogue subsequent to the initial review and requests revisions or explanations that can result in significant delay to protocol approval. By altering the model to one in which the ethical reasoning is included in the original submission, dialogue between the sponsor/investigator and the ethics committee can be initiated upfront. The process would therefore become more efficient and ethics issues would be addressed proactively, directly and more completely.

There are limitations of the Ethics Tool Kit. There may be clinical trial questions that do not fit neatly into the framework we have developed, or the Ethics Tool Kit may be of limited utility when certain methodologies are included in clinical trials, particularly as those methodologies develop and change. For instance, adaptive clinical trials introduce the ethical dilemma of whether and when the investigator should disclose the results to date to prospective participants. If results are disclosed, later prospective participants may not wish to be randomised to what appears, with time, to be the inferior arm. Since this is an emerging issue and there is no international guidance on this dilemma, the working group did not address it in the Ethics Tool Kit.

The 11 Essential Elements are considered a starting point for protocol ethics discussion. Emerging concerns (eg, data transparency, publication policy, recruitment feasibility, innovative trial design) may result in future modifications. Feedback is being actively sought by the MRCT Center, so that the Ethics Tool Kit can continue to be refined and updated. Based on initial online use metrics, it appears that the Ethics Tool Kit is providing a needed educational resource for those seeking guidance on ethical protocol writing. An update is envisioned in 2017, and in early 2017, a survey will be deployed to uniformly collect user feedback on the value of the Ethics Tool Kit.

CONCLUSION

▸ Substantive discussion of specific ethical issues is rarely included in clinical trial protocols.
▸ A total of 11 ‘Essential Elements’ have been identified that should be considered and addressed as appropriate in a clinical trial protocol.
▸ The Protocol Ethics Tool Kit has been developed to support protocol writers, study teams, sponsors, ethics committees and reviewers.
▸ Use of this tool could result in more efficient development and review of clinical trial protocols and may result in wider appreciation of the ethical challenges in clinical research.

1An online feedback discussion forum is available at https://bioethicsresearchreview.tghn.org/community/groups/group/essential_elements/ that captures comments in real time.

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Contributors The paper originated from the work of a multi-stakeholder group convened by the Multi-Regional Clinical Trials Center of Brigham and Women’s Hospital and Harvard (MRCT Center) to develop ethical principles for writers and reviewers of protocols for clinical trials that involve emerging economies. Participants in the working group were self-selected based on relevant expertise and were self-funded. The MRCT Center is supported by voluntary contributions from a variety of entities as well as grants (see http://mrctcenter.org/about-mrct/funding-and-support/). Each of the authors participated in convened meetings that formed the basis of the content, and contributed to writing one or more sections of the toolkit described in the article; each author reviewed a draft of and the final submitted manuscript. The manuscript itself was written and revised by the first two and last two authors listed. The guarantor is Barbara E. Bierer, MD Professor of Medicine, Harvard Medical School and Faculty Co-chair of the MRCT Center; she retains final responsibility for the content.

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REFERENCES


20 Sofaer N, Strech D. Reasons why post-trial access to trial drugs should, or need not be ensured to research participants: a systematic review. *Public Health Ethics* 2011;4:160–84.


Correction


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