At What Level of Collective Equipoise Does a Randomized Clinical Trial Become Ethical for the Members of Institutional Review Board/Ethical Committees?

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ABSTRACT

Background: The conduct of a randomized controlled trial (RCT) is deemed ethical only if we are in state of “equipoise” as to which treatment would be most beneficial for the patients. Individual equipoise applies to an individual clinician or a member of ethical, institutional review board (IRB), whilst collective equipoise refers to the profession as a whole. It is argued that physicians are not bound by the equipoise but their actions are directed by the confines of the expert opinion. Experts can agree or disagree in various proportions on the merit of a given treatment. Hence, the collective equipoise will be often incomplete. In turn, the opinions of content expert in the field expressed as proportion of experts favors one treatment over another (median: 80%; third quartile: 80%). Similarly, half of participating IRB members would approve the study when the median distribution of equipoise among experts was 70% for treatment for leukemia in dogs and 85% for leukemia in rats (and 25% of IRB members would approve such a study even if 100% of experts favors one treatment over another). None of the demographic features of respondents affected collective equipoise.

Conclusions: This is the first study assessing collective equipoise among ethical committee/IRB members. Our study findings show that IRB members perceived that conduct of a trial enrolling humans is unethical when the equipoise level is beyond 80% (80:20 distribution of uncertainty). IRB members require a higher level of equipoise when it comes to testing a new drug in humans than in animals. A relatively high level of equipoise is needed for IRB members to be comfortable to approve trials involving life-threatening situations, children and elderly patients.

Key words: randomized controlled trial, ethics, Ethical committees.

1. INTRODUCTION

There are three key approaches to conducting scientific research in humans: goal oriented, duty oriented and the right-based approaches. The goal oriented approach also termed as utilitarian approach ensures scientifically robust results which promise maximum benefit to the greatest number of future participants and to research in humans (4-6). That is, research should be conducted with a goal to resolve existing uncertainties e.g. to compare the efficacy of antibiotics use vs. watchful waiting for the management of upper respiratory tract infections among children. If there are no uncertainties, there would be no need for clinical research to inform decision-making (7).
Randomized controlled trials (RCTs) are considered as the gold standard for informing treatment decisions as RCTs are based on a deductive method. That is, if the assumptions of the test are met, a positive result obtained via a RCT implies the appropriate causal conclusion. Hence the majority of researchers and participating patients accept RCTs as a tool to resolve existing uncertainties (8) (9). However, the conduct of a RCT is deemed ethical only if we are uncertain that is in state of “equipoise” as to which treatment would be most beneficial for the patients. If we are truly in equipoise, a clinical trial, and particularly a RCT, serves the patients’ interest best, since he or she does not lose out prospectively on the benefits for participation as the treatment, that is being tested, has equal chances to be as beneficial as harmful (6).

Any new study is designed based on the existing knowledge of the treatments being tested. Existing information can range from simply not knowing to having different degrees of uncertainties including a state of equipoise when we are equally positioned in our beliefs between the benefits and harms of a certain treatment or the choice between two or more competing treatments (6, 10). The state of equipoise can be held by researchers, patients and members of the community at large including the research regulatory officials such as members of the institutional review board (IRB) who approve the trials.

Individual equipoise applies to an individual clinician or IRB member, while collective equipoise refers to the profession as a whole (10). Freedman et al argued that physicians are not bound by the equipoise but their actions are directed by the confines of the expert opinion (11). Experts can agree or disagree in various proportions on the merit of a given proposal, and in their beliefs, say if treatment A is superior to treatment B. Hence, the collective equipoise will be often incomplete. In turn, the opinions of content expert in the field (e.g. oncologists) of the proposed trial influence the individuals involved in the conduct of research in humans including IRB members’ decision regarding trial approval. Hence, it is worthwhile to investigate under which circumstances IRB members will be equally split (50:50) in their decision to approve a proposed RCT as a function of the proportion of experts favoring one treatment over another? Accordingly, we conducted a survey of IRB members to assess the degree of collective equipoise necessary for a specific type of trial to be deemed ethical and hence approved by IRB.

2. METHODS

Data collection: We conducted a survey of IRB members at University of South Florida and the IRB members attending the bioethics conference organized in Clearwater, Florida, USA. This paper represents a full description of the previously reported analysis in the abstract form only (12). The survey was distributed to 218 IRB members. The survey was made available as hard copy (paper based) and included six hypothetical scenarios outlining clinical trials targeted at measuring the collective equipoise. We defined the collective equipoise as the situation when survey participants were equally split (50:50) in their decision regarding whether a proposed clinical trial would be ethical to conduct. In each of the five scenarios the choice of the treatment was determined by chance alone (flip of a coin). The opinion of 100 experts in the field expressed as proportion of experts favoring treatment A vs. B in each of the five scenarios was made available to the participants. IRB members were requested to answer the question: “under which circumstances of the experts’ agreement/disagreement would you approve a randomized trial for each of the scenarios listed above?” The survey was modeled after the one described by who assessed collective equipoise in lay people (13). We collected data on demographics including years of service with IRB from the participants. Our study was approved by University of South Florida IRB.

Data analysis: We conducted descriptive analysis including median and interquartile range of approval of trial by participants for the six scenarios. We assessed the difference between the collective equipoise estimates of non-MD IRB members and IRB members who were physicians by employing the Mann-Whitney test (14). We conducted simple logistic regression to determine the impact of demographic variables on the collective equipoise.

3. RESULTS

The response rate of our survey was 33% (71/218). Sixty seven percent (146/218) of the participants were females and 33% (72/218) were males. Fifty eight percent (126/218) were active members of the IRB at the time of the survey. Fourteen percent (31/218) were physicians (holding a MD degree). The median age of participants was 48 years (range: 24-75 years). On average the participants served for 3 years on IRB (median: 3 years, range: 0.5-12 years).

Fifty percent of the IRB members would approve an RCT addressing the efficacy of two drugs for the management of headache even if 80% of experts favor one treatment over another (median: 80%; third quartile: 80%) (Table). Even if 70% (median: 70%; third quartile: 80%) of experts favor one treatment over another 50% of IRB members would approve an RCT addressing the efficacy of two drugs for the management of leukemia. That is, 25% of IRB members would accept enrollment into a trial addressing safety and efficacy of two treatments for the management of leukemia even if 80% of the experts favor one treatment over another (Table). Even if 60% (median: 60%; third quartile: 70%) of experts favor one treatment over another 50% of the IRB members would approve an RCT addressing the safety and efficacy of two antibiotics in the treatment of pneumonia among elderly patients. That is, 25% of the IRB members would accept enrollment into trial assessing safety and efficacy of antibiotics for the treatment of pneumonia among elderly patients even if 70% of the experts favor one treatment over another (Table).
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...treatment over another 50% of the IRB members would approve an RCT addressing the safety and efficacy of two antibiotics in the treatment of pneumonia among newborns recovering from surgery. That is, 25% of the IRB members would accept enrollment into trial testing safety and efficacy of a new drug in humans and children who are considered to be potentially vulnerable to violation of free consent to participate in a trial.

4. DISCUSSION

This is the first survey assessing collective equipoise among ethical committee/IRB members. Findings of our survey indicate that IRB members require a higher level of equipoise (i.e., more uncertainty) when it comes to enrolling a new drug in humans than in animals. The trial enrolling humans most likely to be tolerated is the comparison of analgesics for the management of headache. IRB members responded with a significant variation in their acceptance ranging from 50% to 100%. A relatively high level of equipoise is needed for IRB members to be comfortable to approve a trial involving life-threatening situations, such as trial addressing efficacy and harms of two competing interventions for leukemia. Our survey results show that the highest level of equipoise is required when new drugs are tested in older populations and children who are considered to be potentially vulnerable to violation of free consent to participate in a trial.

Interestingly, under the assumptions that experimental treatment is >80% successful most rational decision for patients themselves is to trust researchers (and by extension IRB members) despite the possibility that the researchers or IRB members may decide to approve the study based on the factors other than patients’ benefits: the likelihood of obtaining successful treatment appears to justify putting oneself in a vulnerable position.

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