Data and Safety Monitoring in Pragmatic Clinical Trials

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Overview

• The need for DSMBs (Jeremy Sugarman)
• Special considerations for DSMBs in PCTs (Susan Ellenberg)
• Challenges for investigators (Greg Simon)
• Discussion
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Background

• Blinding and randomization are powerful research techniques used to minimize bias
• However, emerging experiences and data can pose ethical quandaries for investigators in meeting their obligations to minimize risk to participants
Why have an external approach to monitoring?

- To ensure that participants are not exposed to undue risk
- To ensure that trial will yield usable results
- To balance the interests of patients within the trial with those outside the trial
- To guard trial integrity
What is a Data and Safety Monitoring Board?

• An independent group charged with reviewing the progress, conduct and outcomes of an ongoing clinical trial
Who is on a DSMB?

• Not set in stone
• Clinical experts
• Biostatistician/trialist
• Ethicist? Patient advocate? Investigators? Representative of sponsor?
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DSMBs for pragmatic trials

• What are the special issues for DSMBs for pragmatic clinical trials?
• (ARE there any special issues for DSMBs for pragmatic trials?)
Issues for discussion

• Need for a DSMB
• What a DSMB will monitor
• Participant follow-up
• Data analysis
• DSMB composition
Issue 1: Do PCTs need DSMBs?

• All clinical trials require some monitoring of interim data
• General guidelines for requiring a DSMB apply to pragmatic trials
  • Trials in which participant safety requires regular review of comparative safety and efficacy data
  • Trials intended to have substantial public health impact
• Since pragmatic trials will typically be addressing questions intended to impact health practices, an expert oversight group will be important for most PCTs
Issue 2: what gets monitored?

• Traditional trials: monitor data on safety, efficacy, and quality of study conduct
• These are important in pragmatic trials also
• Possible special issues in pragmatic trials
  • Study outcomes
  • Protocol adherence
  • Eligibility
  • Design factor in cluster randomized trials
What gets monitored: study outcomes

• Study outcomes
  • PCTs may be more likely to include subjective outcomes as primary or key secondary endpoints
  • PCTs may be less likely to incorporate central adjudication of outcomes
  • DSMBs will have to recognize that data may be more variable than in more restrictively designed trials
What gets monitored: adherence

- Protocol adherence
  - A basic tenet of PCTs is to evaluate treatments as they would be given in practice
  - To some, this means no great effort to promote, or even monitor, adherence to protocol
  - DSMBs typically consider monitoring study quality as one of its mandates; may be uncomfortable making recommendations based on observed treatment effects without any sense of how effectively interventions are being administered
  - If adherence is very poor and there is no apparent treatment difference, 2 possibilities
    - Treatments produce similar effects
    - Protocol not followed by investigators and participants
  - If you don’t know anything about adherence, may not be able to conclude anything about relative treatment effects
DSMBs and protocol adherence

- Should a DSMB ignore data on protocol adherence in a PCT? Should these data not even be reported?
- Poor adherence could lead to safety issues in some studies
- Important to distinguish between
  - Lack of adherence as reflecting how a treatment would be used in practice
  - Lack of adherence as reflecting insufficient understanding of trial on part of investigators and/or participants
- DSMBs need to pay some attention to this issue
- May be particularly important to review adherence data by site, to assess need for re-training
What gets monitored: cluster-randomized trials

- For cluster-randomized trials, design often used in pragmatic trials, also important to monitor the “design factor”
  - Intra-cluster correlation coefficient (ICC)—the extent to which results within a cluster will be more similar than results across clusters—is a component of sample size calculation
  - Typically, hard to estimate ICC from prior data—estimates used to design trial may be way off
  - Interim estimates of ICC important to see whether study will have expected power
Issue 3: Participant follow-up

- Pragmatic approaches to follow-up may create challenges for DSMBs.
- Follow-up information will likely be derived from electronic health records (EHRs) in some trials which may be updated on different schedules if different systems are used.
- Follow-up frequency may vary by institution according to local policies.
- Interim comparisons will be more difficult without standardized follow-up schedules.
Issue 4: Data analysis

- Analytical issues
  - Cluster randomization
  - Decentralized analysis
- Philosophical issues
  - Early termination criteria
Data analysis

• Use of cluster designs
  • Many PCTs currently underway with NIH collaboratory or PCORI funding randomize clusters rather than units
  • Analysis of such trials requires accounting for intra-cluster correlation
  • Differing practices among clusters will have to be accounted for in interim analyses
    • Example: minimally restricting usual practice may mean patients in different clusters are followed on different schedules
Potential analytical issue

• Need for de-centralized analysis
  • Privacy concerns may preclude merging data from multiple EHR systems at a central site
  • In such cases, interim analyses may need to be done separately for each site, with summary data only delivered to central statistical group
  • Such arrangements will raise challenges in terms of timeliness of data, quality control and assurance that all analyses have been conducted in identical manner
Interim monitoring strategy

• Early termination for efficacy
  • Since PCTs will be designed to influence practice, could be argued that early termination criteria should be extremely stringent (or maybe not even considered)
  • Will be important to ensure that DSMB and trial leadership are in agreement on criteria

• Early termination for futility
  • When studies compare two “standard-of-care” regimens, questionable whether early stopping for futility should be considered at all
  • As with efficacy, DSMBs and trial leadership must have common understanding of criteria for early termination

• Early termination for safety
Issue 5: DSMB composition

• Clinical and statistical expertise needed
• Will probably be more common to include patient representative
  • PCORI-funded studies require patient partners as members of research teams
  • Studies aimed at questions intended to influence clinical practice may particularly benefit from patient insights
• Expertise in medical informatics may be desirable for some PCTs
  • Use of electronic health data
  • Complex database linkages
  • Natural language processing
The DSMB Charter

• The charter is essentially an agreement between the DSMB, the trial sponsor and the trial investigators about the responsibilities and operation of the DSMB

• The charter will address issues such as
  • Meeting format and frequency
  • Conflicts of interest
  • Statistical approach to monitoring
  • Preparation of meeting minutes

• A Charter is not a formal contract; it guides DSMB actions but the DSMB must be free to exercise its judgment
Greg Simon, MD, MPH

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What’s different in pragmatic trials

- Variable fidelity of or adherence to interventions
- Inference regarding adverse events
- Limited access to outcome data
- “Actionability” of interim analyses
Monitoring intervention fidelity or adherence

• The question: Will this trial support valid inference regarding the benefits and risks of the intervention(s) being tested?
• In a traditional clinical trial, gaps in fidelity or adherence are threats to validity
• In a pragmatic trial, gaps in fidelity or adherence are signal rather than noise
• BUT....Is there any limit to that?
SPOT study example

- Outpatients reporting frequent suicidal ideation on routine depression questionnaires randomly assigned to outreach programs or continued usual care (no contact)
- Outreach invitation via online messaging in EHR patient portal
- Up to three cycles of outreach – patients free to decline or ignore invitation
- Analysis by initial treatment assignment, regardless of intervention participation
- Pilot studies found that 40-45% actively “accepted” invitation to program
- BUT invitations themselves have “active ingredients” of proven interventions
- AND we don’t know how participation is related to actual risk of suicide attempt
Is there a lower bound to adherence? It depends...

- Can we define and measure exposure (or non-exposure) to the intervention?
- What proportion of participants would need to be exposed to detect benefit?
- What do we know about beneficial or adverse selection into participation?
Monitoring adverse events

- The traditional question: Do adverse events signal some risk or potential harm of study intervention(s)?
- The traditional process:
  - Review of individual events (especially unexpected events) for “relatedness”
  - Comparison of event rates for serious adverse events (SAEs)
- In a pragmatic trial:
  - Study teams may not have access to clinical data to assess “relatedness”
  - For treatments in widespread use, a “signal” of harm may be only noise
  - SAEs may be indistinguishable from study outcomes
SPOT study example

- Suicide attempts and suicide deaths would usually be considered SAEs requiring immediate review.
- BUT:
  - those are the study outcomes
  - we are expecting to observe about 700 suicide attempts and 70 suicide deaths
  - they may be ascertained by nothing but an ICD10 code
- SO how would we assess whether suicide attempts or deaths are related to study intervention(s) except by finishing the trial?
Should we monitor adverse events? It depends...

- Unlikely that any “unexpected” event would signal a previously unrecognized risk?
- And comparison of event rates often overlaps with interim analyses (more later)
- But there still may be important questions regarding conflict of interest:
  - “Adverse events” may signal a risk that’s important, whether or not it’s “related”
  - Investigators and study staff must place duty to participants over duty to protocol
  - So monitoring may be indicated – but it’s about a different question
Limited access to outcome data

- Traditional trials rely on data collected and recorded by study staff, so:
  - The study protocol can dictate content and process of data collection
  - Data are available (almost) immediately
  - Study staff control the chain of custody

- Pragmatic trials often rely on the “data exhaust” of health care operations, so:
  - Data collection is controlled (or not controlled) by clinical and business needs
  - Access to data may be delayed by weeks or months
  - No single chain of custody is possible
SPOT study example

- Suicide attempts ascertained through EHR and insurance claims data
  - Often delayed up to 3 months
  - Clinical information to validate or adjudicate may be limited
- Suicide deaths ascertained through state mortality data
  - May be delayed by 18 months or more!
  - Clinical information to validate or adjudicate will be absent
Living with limited access to outcome data

- In many cases, prompt reporting of deaths is neither possible nor useful
- May affect timing (or even feasibility) of any interim analyses
When can we act on interim analyses?

- In a traditional clinical trial – once the question is answered, we should:
  - Stop assigning research participants to an inferior treatment
  - Advise clinicians and policy-makers regarding new evidence
- In a pragmatic trial
  - Outcome data may accumulate slowly (at different rates from different sources)
  - The threshold for action may be less clear
SPOT study example

• Delayed (and complex) schedule for outcome data:
  • Outcomes accumulate over 12-18 months
  • Greater delay for suicide deaths than non-fatal suicide attempts
  • Should not over-value early over sustained intervention effects

• Threshold for health system action is not clearly established (and will likely depend on cost as well as benefit)

• Consider practical consequences of stopping recruitment or intervention delivery:
  • For evidence of benefit:
    • Patients in participating health systems would no longer be offered effective programs
    • Timing of widespread implementation uncertain
  • For evidence of harm:
    • Patients in participating health systems would no longer be offered harmful programs
    • Widespread implementation would be avoided
Are interim analyses actionable? It depends...

- Consider timeline of data availability:
  - How soon could you detect meaningful or important difference?
  - What biases could be introduced by using incomplete data?
- Consider effects of early termination on potential study participants and others affected by condition of interest
- Consider different thresholds for detecting benefit and harm
DISCUSSION