Choosing a Research Question:
Implications for Efficient Clinical Trials

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Introduction

Choosing a research question as first step in research endeavors

Types of research questions

Process of working from one question to the next – building a research “portfolio”

Choice of research question and implications for design of studies

Influence on study design and sample size

Most efficient use of resources
Boundaries between Research and Practice

Clinical research – designates activity designed to test hypothesis, permit conclusions, and thereby develop or contribute to generalizable knowledge (expressed in theories, principles, and statements of relationships)

Clinical practice – interventions designed solely to enhance well-being of individual patient or client and have reasonable expectation of success, purpose to provide diagnosis, preventive treatment or therapy to particular individuals

Belmont Report on Ethics in Human Experimentation
Scientific and Ethics

Study that cannot contribute to generalizable knowledge is not ethical

Puts patients at risk of harm (even from minor inconvenience) for no benefit to anyone

Scientific validity is not a “nice to have” but a requirement of all research

Validity = ability of study to correctly answer research question posed
Science

“It is not what the man of science believes that distinguishes him, but how and why he believes it. His beliefs are tentative, not dogmatic; they are based on evidence, not on authority or intuition.”

Bertrand Russell, A History of Western Philosophy, p. 527.
Focusing the Question
Choosing a Topic

Idea come from life experiences

Clinical practice

“What do I have” (diagnosis)

“How bad is it” (prognosis)

“Can I give it to my family?” (natural history)

“How did I get this?” (etiology)

“Will this stuff help me?” (treatment or prevention)

Clinical research often focuses on biology but can focus on other areas

“Why do people come to the doctor for disease X?”

“What behaviors influence outcome in disease Y?”
Choosing a Topic

Developing better tools to evaluate and describe disease is a good place to start.

Need to characterize natural history of disease in order to study it.

Need to have valid measures of outcomes in order to 1) measure them and 2) describe risk factors for outcomes.

Example of measurement tools:

- Seriousness/severity of disease (comparing baseline factors with outcomes)
- Developing outcome measures (measures of morbidity e.g. functional status and symptoms)
Descriptive Research

"Classification is fundamental to the quantitative study of any phenomenon. It is recognized as the basis of all scientific generalization and is therefore an essential element in statistical methodology. Uniform definitions and uniform systems of classification are prerequisites in the advancement of scientific knowledge. In the study of illness and death, therefore, a standard classification of disease and injury for statistical purposes is essential."

Areas for Investigation
Many Related Areas Other than Biology
Choose a Broad Topic

Choose an idea that is interesting – you have a lot of work to do!

Choose a topic that is timely and relevant (need to answer the “so what” question) – a question worth answering

Choose a topic that is answerable – keep in mind time and resource constraints


Feasible – BUT feasible and invalid = unethical

Interesting

Novel

Ethical

Relevant

Mentor can help with all these facets of project
Focusing the Question
Focusing the Question

Need to evaluate the medical literature and other sources to evaluate current knowledge in research area

In many areas, much of what we know is less certain than generally believed

Review and guidelines are places to start, but represent synthesis of others views on the data – need to review for yourself
What Do We Know Already?
The “Knowledge Gap”

How do you define the problem – clinical features, signs, symptoms, laboratory values

How do you diagnose the problem? What is impact of diagnosis on outcome?

How large is the problem? Magnitude of problem in terms of number and types of persons affected

What is impact of problem? Attributable morbidity and mortality (often assumed)

What are risk factors for getting problem? (not necessarily causal)
What Do We Know Already?
The “Knowledge Gap”

6. What is prognosis of problem? Morbidity and mortality

7. What factors modify prognosis independent of treatment? (Confounders)

8. What interventions can mitigate problem? Interventions can include drugs, devices, biologics, and behaviors

9. What factors are effect modifiers of treatment? Effect size differs depending on presence of factor

10. How do interventions compare to each other?
What Do We Really Know?

“The greatest obstacle to discovery is not ignorance, it is the illusion of knowledge”

- Daniel Boorstin
What Do We Know Already?
The “Knowledge Gap”

What do we know about previous questions?

Often challenging to convince others we don’t really know what we think we know

Argumentum ad verecundiam fallacy (argument from authority)

What is quality of data? Validity, reliability and precision, biases in previous data

Has evidence been independently confirmed?

Is evidence consistent across different populations? Is a difference biologically plausible?

For interventions: dose, duration of therapy, combination therapy
Overall Research Plan

After evaluating what is known, need to focus on one part of research gap, one piece of the puzzle

How does this research fit into an overall plan? What happens after this study (by you or someone else)?

A single study cannot possibly answer all questions about a topic

Research fits into an overall model/theory of a problem
Feasibility

What kinds of information do you need to answer the question?

Population – both test and control groups in analytical study

Exposures

Outcomes

What kinds of information are available?

What resources are needed to obtain data needed?

Is there access to resources needed?

Feasibility does NOT mean using invalid methods because that is “the best that can be done”
Focusing the Question
Developing Hypothesis or Description

Hypothesis – a statement about what investigator believes to be true about nature and relationships of two or more variable to each other

Hypothesis testing entails a comparison, but not all research is comparative

Differentiate qualitative and quantitative research

Differentiate descriptive from analytical research
Developing Hypotheses
Qualitative and Quantitative Research

Qualitative

Aim is complete detailed description – words

Develop observations for further testing

Only know roughly in advance what to look for

Early phase of research project

Design emerges as study unfolds

Researcher is data instrument

Quantitative

Aim to classify features, count them - numbers

Construct statistical models to explain observations

Clearly state in advance what to look for

Later phases of research project

All aspects carefully designed before data collection

Researcher uses tools (questionnaires, equipment) to collect numerical data
Developing Hypotheses
Descriptive and Analytical Research

Descriptive research – provide an account and delineate components of a problem

Case report

Case series – more data does not necessarily increase validity

Analytical research – testing one of more hypotheses in a quantitative fashion

Distinction not as clear as descriptive research often contains comparisons (but cannot assess causality) and analytical research often contains descriptions

Push for “hypothesis driven” research tends to make descriptions sound less valuable but descriptions help form hypotheses
Choosing A Design
Types of Clinical Studies
Examples of Hypotheses

The more specific the better:

“Antibiotics are effective in acute otitis media in children”

“Amoxicillin is effective in acute otitis media in children who are between 2 and 6 years of age”

“Amoxicillin is effective compared to placebo in reducing pain in children ages 2 to 6 years with initial episodes of acute otitis media”
Focusing the Question
Specific Aims and Objectives

Choosing an overall research questions gives you a “why” (the rationale for doing the study)

Next need to answer the questions related to specific measurements and define them:

Who – define population under study

Where – setting in which study will occur

When –

what time frame of analysis; “from January 2000 to January 2009”

Prospective or retrospective – hypothesis in relation to data, not how collected

What – variables of exposure, intervention and outcome (content validity)

How – what tools to use to measure variables (construct and criterion validity)

Need to be as specific as possible regarding measurement variables and how to measure them

Failing to plan is planning to fail

Avoid circular or vague language: “Clinical outcomes will be divides into clinical success and failure”
Right Tools for the Job
Common Pitfalls

Letting “feasibility” issues change the question to one not worth answering or answering it in an invalid way

Taking on too many questions and thereby answering none

Lack of clarity of hypothesis and lack of clarity on study design

Vague specific aims and variables and unclear measurement properties of tools
Designing Trials Efficiently
Definitions

Clinical trial – a controlled prospective study enrolling human subjects often used to evaluate the effectiveness and/or harms of interventions in treatment, prevention or diagnosis of disease

Efficiency –

(in physics) ratio of useful work to the energy supplied to it

In clinical trials, getting valid and reliable answers to important questions with the least amount of resources

Does not mean putting patients at risk because of less valid data

Lower sample size does not mean less work = MORE planning
Lower Sample Size = More Planning

“Clinical trials with small numbers of participants, however, must address broad sets of issues different from those that must be addressed in trials with large numbers of participants. It is in those circumstances of trials with small sample sizes that approaches to optimization of the study design and data interpretation pose greater challenges.”

Small Clinical Trials: Issues and Challenges, Institute of Medicine, 2001
http://www.nap.edu/catalog/10078.html p. ix
Definitions

What is a “large” or “small” clinical trial?

IOM defines a “large” trial as one that has adequate sample size to answer the primary research question = “large enough”

A trial with very few participants may still have adequate statistical power e.g. if effect size is large

Balance between exposing research subjects to potential harms of experimental interventions with obtaining valid answers
Underpowered Studies and Ethics
Underpowered Studies and Ethics

“A proposed study that cannot answer the question being asked because the necessary sample size cannot be attained should not be conducted on ethical grounds. That is, it is unacceptable to expose patients or research participants to harms, even inconveniences, if there is no prospect that useful and potentially generalizable information will result from the study.”
Small Clinical Trials – Last Resort

“The importance of conducting small clinical trials only when there are no alternatives cannot be overemphasized. The committee is not encouraging the use of small clinical trials, but, rather provides advice on strategies that should be considered in the design and analysis of small clinical trials when the opportunity to perform a randomized clinical trial with adequate statistical power is not possible. In doing so, it recognizes that small clinical trials frequently need to be viewed as part of a continuing process of data collection. Thus, for some trials it might be impossible to definitively answer a research question with a high degree of confidence. In those cases, perhaps the best that one can do is assess the next set of questions to be asked.”
Concerns About Small Clinical Trials

Small numbers increase variability and leave much to chance

Statistically significant outcomes may not be generalizable (only apply to circumstances in trial)

Too many variables to assess cause and effect

Only able to discern gross effects and limited ability to analyze covariates

Incapable of identifying adverse events

Small Clinical Trials: Issues and Challenges, Institute of Medicine, 2001
http://www.nap.edu/catalog/10078.html p. 15
Situations Where Smaller Clinical Trials Justifiable

Rare diseases

Unique study populations (e.g. astronauts)

Individually tailored therapies

Environments that are isolated

Emergency situations

Public health urgencies

Small Clinical Trials: Issues and Challenges, Institute of Medicine, 2001
Small vs Efficient

While small clinical trials are a last resort, efficient clinical trials are always justifiable

Different methods to improve efficiency are useful (or not) depending on disease under study and research question/setting
Components of Clinical Studies

1. Clear objective of study

2. If comparative (rather than descriptive) quantitative comparison with control group

3. Select patients for inclusion in study

4. If comparative, baseline comparability of groups compared

5. Minimizing bias of study

6. Well-defined and reliable outcome measures (patient-centered)

7. Appropriate statistical analysis
Review of Sample Size Considerations
Normal Distribution of a Sample
Sample Size – Superiority Trials

Clinical trials compare average effects in groups of subjects administered intervention to those not administered intervention

Examine if populations differ by more than chance (for superiority trials)

Example: Two groups with point estimate for means of 50 and zero, sample size of 12 per group, SD = 60
Sample Size – Superiority Trials

In this example

Blue area represents power (in this case 0.497)

“critical region” of 0.05 test represented by dashed lines and red area

To show difference due to greater than chance, want mean of one curve to be outside of red area
Sample Size

Sample size in this example is 24 per group

As sample size increases, overlap between curves decreases (assuming there really is a difference to show)

Blue area increases = power is 0.80

Mean of one group now outside of “critical area”

Notice still a good deal of overlap – only mean value is outside critical area
Sample Size

Sample size increased to 48 per group

Distribution of data narrower and more precise

Power = 0.98

Mean well outside of “critical area”
Sample Size

Want to select sample size large enough to show a difference if there is one to detect, but not too large.

Do not want to expose subjects unnecessarily to harm since this is an experiment evaluating interventions with unknown harm/benefits.

Use of resources - time, effort and money.

Sample size based on four parameters (“ingredients”)

Type 1 error- usually specified as 0.05 two sided (0.025 on either side of curve)

Type 2 error (1-type 2 error is power) usually specified as 0.10 to 0.20 (power of 80%-90%)

Standard deviation of data (variability)

Treatment difference – Difference between point estimate of effect for intervention and point estimate for effect with control (delta)
Pictorial Representation of Sample Size
Pictorial Representation of Sample Size
Efficient Clinical Trials

Ways to decrease sample size

Focused and relevant research question

Changing error rates (not suggested)

Enhancing effect sizes

More homogenous populations

Choosing populations in whom effect size is larger

Optimizing exposure

Continuous instead of dichotomous outcomes

Selection of outcomes

Decreasing variability

More sensitive/specific measures

Assuring follow-up of enrolled subjects

Study designs

Cross-over

N=1 studies

Sequential trial designs (e.g. dose-escalation studies)
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1. Focusing the Research Question

Research question needs to be one worth answering and of public health importance

Need to focus question – more questions mean greater sample size or less clear answers = simplify

“Many trials include measurements to try to figure out why the trial didn’t work after it has failed” – the post-mortem on “what the experiment died of”
1. Focusing the Research Question

Sample size is calculated AFTER one decides on a research question

Starting out with a sample size and working back to “what can I get for this” is not justifiable in terms of choosing unrealistic or clinically meaningless/unachievable effect sizes
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2. Changing Error Rates

Error rates of type 1 = 0.05 and type 2 = 0.10 or 0.20 are by convention

But...false positive error rate of 1 in 20 trials is actually a low level of evidence; need justification to deviate from this

Increasing type 2 error rate increases likelihood of false negative conclusions = spending resources for unclear answers

Error rates measure random error (by chance) but not bias due to poorly designed study
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Enhancing Effect Sizes

Trials measure average effects in groups of subjects – “one size fits all” approach may not be correct

More homogenous populations can both decrease variability and increase effect sizes in presence of effect modification

Effect modification is presence of quantitatively different effect sizes of an intervention based on a baseline variable (e.g. drug is more effective in older people vs younger people)

Requires knowledge of natural history of disease and evidence from prior trials

Choosing population in which effect size is larger decreases sample size
Examples

Enrolling subjects in trial in whom effect is expected to be zero

Dilutes effect size

Ethical issues of exposure to harm for no benefit

Trastuzumab (Herceptin) in breast cancer

Mechanism of action by binding to HER2 proteins in person with specific genetic mutation

20% to 30% of person with breast cancer have this mutation

Potentially harmful in those without the mutation
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Optimizing Exposure

Many trials include only one dose of an intervention with wide inter-individual variability in exposure

Optimize dose based on pre-clinical and early clinical studies – pharmacokinetics and pharmacodynamics

Forms a hypothesis to test

Not a substitute for clinical trials

Standardize exposure of interventions - need unblinded third party to do this to maintain blinding in trial
Efficient Clinical Trials

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Continuous vs Dichotomous Outcomes

Continuous outcomes have more power to detect differences since uses all the data

Dichotomous outcomes require

1) categorization – assumes all data in a single category are similarly important which may not be true
2) choosing correct time point to evaluate – if you’re wrong, you miss it

Requires more frequent data capture – patient diaries or phone contact collected in systematic way
Time to Event - Cholera
Efficient Clinical Trials
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Sequential trial designs (e.g. dose-escalation studies)
Common or Composite Outcomes

Choosing more common events or composite outcomes increases number of events and increases power to detect differences

Some problems with interpretation:

Only use when outcomes measured are of similar importance to patients – if driven by less important outcomes may mask inferiority on more important outcomes

Does not necessarily imply beneficial effect on all part of a composite
Composite Endpoints - Issues
Efficient Clinical Trials

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More Sensitive/Specific Measures

All measurements composed of the true value plus some associated error

Error can be of two types:

Random – by chance alone

Systematic bias – based on individual biases and interpretations

Decreasing error (“noise”) in relation to true measure (“signal”) allows smaller sample size
Non-Standardized Measures and Error
Non-Standardized Measures and Error


16 oncologists asked to measure 12 simulated tumor masses, two pairs of which were identical in size

Allowed assessment of 64 measurements by same investigator and 1920 comparisons by different investigators

25% “reduction” as “response” for identical size masses = 19% response rate by same investigator, 25% between investigators (measurement error alone)
Surrogate Endpoints

Researchers often suggest biomarkers as “surrogate endpoints” in clinical trials – NOT direct measures of patient benefit

Idea originally was to decrease follow-up time in chronic diseases – keep following subjects to validate biomarker (e.g. viral load in HIV/AIDS, cholesterol in stroke/MI prevention)

Why use a surrogate in an acute disease when one can measure actual clinical outcomes?

Surrogate as part of composite outcomes drive the entire outcome since more common
Strengths and Limitations
Efficient Clinical Trials

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Follow-Up and Missing Data

Enrolling subjects and then losing them to follow-up or missing data (failure to collect or losing it) results in effort for no gain

Requires planning on part of researchers and work during the trial to make it as easy as possible for research subjects to return

Phone calls and reminders

Transportation

Home visits

Subjects who don’t follow protocol are not “missing” and should be included in Intent to Treat (ITT) analyses
Efficient Clinical Trials

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Study Designs

Certain kinds of study designs can decrease variability and thereby decrease sample size.

Cross-over and n=1 trials both use subjects as their own controls.

Randomize subjects to receive one intervention or the other (sometimes with wash out period in between).
Study Designs

Limitations of cross-over and n=1 trials

Most useful in chronic illnesses with stable course of disease

If effects carried over from one period treatment to the next then can bias study results

“Period effect” for instance in seasonal diseases

Most useful in diseases where treatment effect is rapid onset or rapid cessation when intervention stopped
Study Designs

Sequential trial designs often used in Phase 1 dose escalation clinical trials

Based on pre-defining what level of adverse events or not will allow progression to the next dose

Makes decision based on data acquired during trial
Study Designs

“Adaptive” designs - word that describes a variety of changes in trial design based on data accumulated during the trial

Sequential trial dose escalation design is one form but more challenging when modifying other variables like the outcome measure

Advantage = more subjects assigned to more successful treatment

Disadvantage = heterogeneity of subjects based on important risk factors which change as trial progresses introduces bias over time
A caveat…

All the modification we have discussed apply to SUPERIORITY trials

Non-inferiority is misnomer – does not mean “not inferior” as to show an intervention is “equal” or “not inferior by any amount requires showing superiority

Biases which trend results toward no difference in a superiority trial (like too small a sample size) result in false positive conclusions in non-inferiority trials

Non-inferiority does not answer question of added benefit of new interventions; use only in selected situations (e.g improved convenience)
Conclusions

Developing an efficient trial starts with planning and a good research question

Question comes first, sample size second

Various methods to increase effect sizes and decrease variability, when applied in the correct setting, can provide valid and reliable answers to important public health questions

For some diseases, developing the tools (better outcome measures, better data on natural history) is a good start to better trials