Statistical Considerations for Early Phase and Biomarker Studies

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Outline

- Overview
- Dose-finding trials
- Safety and efficacy trials
- Statistical and design considerations
- Correlative biomarker studies
Overview
Clinical trial definition

A clinical trial is an experiment testing a medical treatment on human subjects.

- Not all human research studies are experimental.
- The experiment is what distinguishes clinical trials from other forms of medical research.
- What do we mean by experiment?

“...the essential characteristic that distinguishes experimental from non-experimental studies is whether or not the scientist controls or manipulates the treatment ... under investigation.”

(Piantadosi, Clinical Trials: A Methodologic Perspective, 2005)
Equipoise

- Scientific uncertainty about the superiority of one treatment versus alternative
- Ethical imperative that study participants not be disadvantaged
- Supports comparative trial design to resolve uncertainty
- If consensus about superiority exists, trial is unethical

*From the original* “... at the start of the trial, there must be a state of clinical equipoise regarding the merits of regimens to be tested, and the trial must be designed in such a way as to make it reasonable to expect that, if it is successfully conducted, clinical equipoise will be disturbed.” (Freedman, 1987)
Stages of trial design in drug development

- **Dose-finding trials**
  - Phase I
  - Designed to find the best safe dose
  - Traditionally $\leq 30$ patients

- **Safety and efficacy trials**
  - Phase II
  - Efficacy signal
  - Traditionally single arm studies with 20 - 80 patients
  - Recent increase in randomized studies for targeted therapies

- **Comparative trials**
  - Phase III
  - Definitive trial against standard of care
  - Multi-center
  - Hundreds to thousands of patients
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Dose-finding trials
Phase I trial goals

- **Classic Phase I trials**
  - Cytotoxic agents - “more is better”
  - Find the highest dose that is deemed safe = maximum tolerated dose (MTD)
  - Determine recommended phase II dose (RP2D)
  - Monitor dose-limiting toxicities (DLTs)
  - MTD = highest dose with DLT rate \( \leq x\% \)
    (usually \( x\% = 20\% - 40\% \))
  - Determine schedule
  - Evaluate safety and toxicity
  - Assess pharmacokinetics

- **Newer Phase I trials**
  - Molecularly targeted agents
  - Find dose considered safe with optimal biologic/immunologic effect
  - Goal is to optimize ‘biomarker’ response within safety constraints
  - Identify ‘target’ population
Classic phase I assumptions

Efficacy and toxicity both increase with dose

![Graph showing the relationship between dose level and probability of DLT and Efficacy](image-url)
New paradigm for molecularly targeted agents

- Agent selective for a molecular ‘target’
- Disrupt carcinogenesis by interfering with specific pathway
- Can be less toxic than traditional cytotoxic agents
- Implications for phase I design
  - Toxicity may be extremely low
  - Need to ensure agent ‘hits’ the target
  - Efficacy may not increase monotonically with dose
  - MTD being replaced by optimal biologic dose or maximum administered dose
Possible relationships for targeted therapies

- Efficacy not necessarily monotonically increasing with dose
- Low toxicity

![Graph showing possible relationships between dose level and probability of DLT and efficacy](image)
Phase I designs

- **Algorithmic**
  - Follows a prescribed set of decision rules for dose-escalation
  - Standard 3+3 design
  - Accelerated titration

- **Model-based**
  - Uses accumulating toxicity data to determine dose for next patient
  - Continual reassessment method (CRM) (or other flavors - mCRM, TITE-CRM)

Classic phase I “3+3” design

**Dose escalation/de-escalation decision rules**

For a pre-specified set of doses (usually between 3 and 10), treat 3 patients at dose-level \( k \)

1. If 0 of 3 patients experience a DLT, escalate to dose \( k + 1 \)
2. If 2 or more of 3 patients experience a DLT, de-escalate to level \( k - 1 \)
3. If 1 of 3 patients experiences a DLT, treat 3 more patients at level \( k \)
   1. If 1 of 6 experiences a DLT, escalate to dose \( k + 1 \)
   2. If 2 or more of 6 experience a DLT, de-escalate to level \( k - 1 \)

- MTD = highest dose at which at most 1 out of 6 patients experiences a DLT
- Therefore, target DLT rate is 33%
- Common to include *expansion cohort* at MTD to obtain additional safety data
Accelerated titration design

- Starts with single patient cohorts
- Approximate doubling of successive doses (96% - 100% dose escalation)
- First DLT or second moderate toxicity in any dose cohort ⇒ expand cohort to 3 patients
- Revert to 3+3 design for subsequent cohorts with 40% dose escalation steps
- Advantages over 3+3
  - Requires fewer patients overall (efficiency)
  - Fewer patients treated at non-efficacious doses
  - Increase precision of RP2D
Continual reassessment method

- Bayesian - uses both accumulating data and prior beliefs to determine actions
- Dose cohorts not specified in advance
- *Learn* best dose assignment for next patient based on toxicity data accumulated at all prior and current dose levels
- Assumes mathematical model relating dose to toxicity
- Requires specification of targeted DLT rate (not 33% by default)
- Patient cohort size must be specified
- Stopping rule - usually based on a pre-specified total sample size
The Bayesian paradigm

- Prior beliefs
- Data
- Posterior inference
- Update prior
CRM dose-toxicity curves
Pros and cons

- **Algorithmic designs ...**
  - Are simple and easy to understand
  - Are efficient - small sample sizes
  - Can be implemented without involvement of a statistician
  - Enjoy wide-spread acceptance (PRCs, IRBs, FDA)

- **Model-based designs ...**
  - Are more challenging to understand
  - Treat more patients at efficacious doses
  - Treat fewer patients at toxic doses
  - Identify the MTD with greater precision
  - Require ongoing involvement of statistician
  - Can be difficult to implement at multiple institutions
Safety and efficacy trials
Phase II safety and efficacy trials

- Provide additional information about toxicity profile
- Provide preliminary information on whether a treatment is efficacious
- Small ⇒ only large treatment effects are detectable
- Quick ⇒ efficacy measured using short-term endpoints

Phase II endpoints

- Surrogates for phase III gold standard, overall survival
- Response is conventional endpoint for cytotoxic agents that cause tumor shrinkage
- Progression free survival (time from enrollment to documented disease progression) is increasingly the endpoint for MTAs
- MTAs may not cause tumor shrinkage, but may prolong time to progression
Conventional phase II designs

- Single arm
  - Efficacy rate in study population compared to historical control rate
  - Control rate may not be well-defined
  - Study population may lack comparability to historical control population

- Randomized selection design
  - “Pick the winner”
  - Patients randomized to two (or more) arms
  - No head-to-head comparison of arms, but rather comparisons to null historical rate
  - Goal is to identify best dose/schedule/regimen to take forward into phase III when there is no *a priori* information that one is preferable

- Randomized with control arm
  - Control arm ensures historical rate “on target”
  - Control arm *not* included for head-to-head comparison (due to small sample size)
Conventional phase II design issues

- Issues with respect to molecularly targeted agents
  - Lack of reliable historical control population ⇒ no comparator historical rate
  - Response inappropriate endpoint when tumor shrinkage not expected
  - Lack of historical data for appropriate endpoint like PFS

- General issues
  - Lack of randomization ⇒ no internal control
  - Treatment-trial confounding (Estey and Thall, Blood 2003; 102:442-8)
  - Unmeasured trial effects (e.g. differences in supportive care, institutional practices, unknown patient characteristics) can not be distinguished from treatment effects
Phase II randomized trials

- Head-to-head comparison
- Increasingly the “norm” for phase II oncology trials
- Requires 2 - 4 times the number of patients as conventional designs
- Increased demand on resources (time and money)
- Standard of care (SOC) + “A” versus SOC (or SOC + placebo)
- Adaptive designs (Gallo et al., *J Biopharm Stat* 2006; 16:275-83)
  - Trial characteristics modified ‘as-you-go’ based on cumulative data
  - Design-based rather than *ad hoc* modifications
  - Trial validity and integrity remains intact
  - Adaptive randomization - increase patients assigned to arm with more efficacious agent
  - Adaptive sample size - re-estimation based on revision of underlying assumptions
Biomarker incorporated designs

- Randomized phase II designs that utilize biomarker status
- Focus on predictive markers
- Often goal is to assess treatment efficacy and validate marker

**Selected designs**
- Enroll patients most likely to benefit - marker positive patients
- Requires biologic accuracy of selected marker
- Reliable marker measurement
- Fails to identify other patient populations who may benefit
- Does not allow assessment of marker-outcome association

**Unselected designs**
- Marker-by-treatment interaction design
- Marker-based strategy design
Marker-by-treatment interaction design

Sargent D J et al. JCO 2005;23:2020-2027
Marker-based strategy design

Sargent D J et al. JCO 2005;23:2020-2027
Statistical and design considerations
Hypothesis testing

- Didactic example - Randomized phase II trial of paclitaxel/carboplatin ± enzastaurin (PCE/PC) in advanced ovarian cancer patients (Vergote et al., *JCO* 2013; 31:3127-3132)

- To compare competing treatments, we begin with rival hypotheses of their relative effect.
  - Null hypothesis - statement of no effect ($H_0$)
  - Alternative hypothesis - statement reflecting conclusion if trial results suggest null is implausible ($H_A$)

$H_0$: Advanced ovarian cancer patients derive no additional clinical benefit from the addition of enzastaurin to paclitaxel/carboplatin relative to those treated with paclitaxel/carboplatin alone.

$H_A$: Advanced ovarian cancer patients derive significant clinical benefit from the addition of enzastaurin to paclitaxel/carboplatin relative to those treated with paclitaxel/carboplatin alone.
Formalizing hypothesis testing

- ‘Clinical benefit’ needs to be articulated in terms of a measurable quantity
- Depends on the trial’s primary endpoint
- If endpoint is response, benefit measured by difference in response rate
- If endpoint is a time-to-event measure, benefit measured by hazard ratio (HR)
Hazard ratio detour

- \( HR = \frac{\text{risk of event in trt grp}}{\text{risk of event in ctl group}} \)
- \( HR > 1 \) indicates increased risk of event in treatment group relative to control
- \( HR < 1 \) indicates decreased risk of event in treatment group relative to control
- \( HR = 1 \) indicates equivalent risk of event in treatment and control groups
- Interpretation
  - \( HR = 0.80 \Rightarrow \) “There is a 20% reduction in the risk of death (progression, recurrence) comparing treated to control patients.”
  - \( HR = 1.20 \Rightarrow \) “There is a 20% increase in the risk of death (progression, recurrence) comparing treated to control patients.”
Back to our hypotheses

If endpoint is response ...

\[ H_0: p_{\text{PCE}} = p_{\text{PC}} \]
\[ H_A: p_{\text{PCE}} > p_{\text{PC}} \]

where \( p_{\text{PCE}} \) is the response rate in the PCE arm and \( p_{\text{PC}} \) is the response rate in the PC arm.

If endpoint is a time-to-event measure

\[ H_0: HR_{\text{PCE}:\text{PC}} = 1 \]
\[ H_A: HR_{\text{PCE}:\text{PC}} < 1 \]
Conducting inference and quantifying evidence

- Based on the evidence in the data, one of two decisions is made
  1. Reject $H_0$ in favor of $H_A$ ⇒ sufficient evidence to rule out the null
  2. Fail to reject $H_0$ ⇒ insufficient evidence to rule out the null
- Evidence is quantified using a $P$-value
- If the null hypothesis is true, it is possible to observe what looks like a treatment effect by pure chance
- $P$-value is the probability of observing a result at least as extreme as what was observed in the current study if the null is true (i.e. if there really is no difference)
- $P$-value quantifies the probability of a spurious finding
- Usually require $p$-value < 0.05 to reject $H_0$ and declare statistical significance
- $p$-value ≥ 0.05 ⇒ fail to reject $H_0$ and conclude we can’t rule out chance as a plausible explanation for any observed difference
Confidence intervals

- Provides range of values quantifying uncertainty associated with an estimated parameter.
- The wider the interval the greater the uncertainty in the estimate.
- The more narrow the interval the greater our faith in the estimate.
- Typically report 95% CI.
- The “95%”-part of the interval means that if you were able to replicate the exact same study an infinite number of times, 95% of the resulting CIs would contain the true parameter of interest (not a particularly practical interpretation).
Putting some of this together ...

In the randomized phase II PCE vs PC trial in ovarian cancer,

“A comparison of ... PFS for the two regimens ... was not statistically significant \((P = 0.367)\). HR at the time of final analysis was 0.80 (95% CI, 0.50 to 1.29).”
Interpretation and conclusions

- Enzastaurin does not significantly increase time to progression when added to a regimen of paclitaxel/carboplatin in advanced ovarian cancer patients ($P = 0.367 \geq 0.05$)

- Adding enzastaurin to paclitaxel/carboplatin therapy conferred a 20% reduction in the risk of progression ($HR = 0.80$), although this finding was not significant. That is to say, although we observed an improvement, we can not rule out chance as a plausible explanation for the reduction in risk.

- Our best estimate of the hazard ratio is 0.80, but the data are also consistent with a HR ranging from 0.50 to 1.29 (95% CI, 0.50 to 1.29). Thus, the data are consistent with equivocal conclusions - a hazard ratio of 0.50 indicates a 50% reduction in the risk of progression while a hazard ratio of 1.29 indicates a 29% increase in the risk of progression.
Actions and errors

Of course it is always possible to make a mistake. The table below details the two types of errors one can commit in conducting a statistical test.

<table>
<thead>
<tr>
<th>Action</th>
<th>$H_0$ true</th>
<th>$H_A$ true</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reject $H_0$</td>
<td>Type I error</td>
<td>No error</td>
</tr>
<tr>
<td>Fail to reject $H_0$</td>
<td>No error</td>
<td>Type II error</td>
</tr>
</tbody>
</table>

- A type I error is the probability of rejecting the null given that the null is true.
- We denote the probability of a type I error as $\alpha$.
- A type II error is the probability of failing to reject the null given that the alternative is true.
- We denote the probability of a type II error as $\beta$. 
The power of a test is the probability of correctly rejecting the null in favor of the alternative. A well-designed study will strike a balance between acceptable levels of type I error (usually 0.05) and power (often set at a minimum of 0.8).

Remember ...

- A type I error is rejecting the null when you shouldn’t
- Power is the probability of rejecting the null when you should
Planned early data looks

- Early stopping for efficacy
  - Is there sufficient evidence to conclude treatment is significantly better?
  - Ethical imperative not to continue to randomize patients to a therapy known to be inferior
  - Inflates type I error rate

- Early stopping for futility
  - No hope of being able to demonstrate treatment efficacy
  - Sufficient data to answer scientific question
  - Unethical to continue to randomize patients in a trial with no additional benefit
  - Inflates type II error rate (decreases power)

- Early stopping for safety
Control of bias and reduction of random error are two major objectives in statistical design considerations.

- **Unbiased**
  - Small random error: Graph A
  - Large random error: Graph B

- **Biased**
  - Small random error: Graph C
  - Large random error: Graph D
Sources of bias

1. *Selection bias*
   - Patients in arms systematically different with respect to prognostic factors
     - Bias the observed treatment effect
     - Can influence internal validity
   - Study cohort not representative
     - Can influence external validity
     - Compromises generalizability

2. *Treatment/procedure selection bias*
   - Healthier patients selected for a particular treatment
   - Systematic difference in composition of treatment groups
   - Can bias treatment difference
3. *Postentry exclusion bias*

- Inappropriate exclusion of eligible and enrolled subjects from the analysis
- Exclusion often due to seemingly reasonable clinical reasons
- Breaks the ‘experimental paradigm’
- Example - subjects that fail to complete therapy are excluded from the analysis
- Example - subjects that die due to ‘other’ causes are excluded from an analysis of overall survival
4. Selective loss of data

- Loss of data resulting from unworkable or suboptimal outcomes or errors in study conduct
- Endpoint poorly selected for patient population
- Frequency of follow-up inappropriate for assessment of desired endpoint or is not followed as specified in protocol
- Example - patient population is seriously ill cohort and endpoint is based on patient self-report; endpoint may suffer from survivor bias
- Example - endpoint is time to progression; follow-up with patients every 6 months may be inadequate for accurate assessment
Sources of bias (cont.)

5. Assessment bias

- Patient self-assessment lacks objectivity
- Clinician assessment can be influenced by expectation of treatment effect
- Can bias endpoint in direction of prior expectation

6. Uncontrolled confounders

- Confounder is a variable that masks the true treatment effect
- Common confounders are age, race, gender, disease severity, comorbidities
- Example - treatment arm is significantly younger than placebo arm, and outcomes in older patients are more severe
Controlling for bias by design - Randomization

- Patients randomly assigned to treatment
- Controls for:
  - Selection bias
  - Treatment bias
  - Uncontrolled confounders
- Types of randomization
  - Simple
  - Permuted block
  - Stratified permuted block
  - Adaptive
  - Group
- Allocation ratio (1:1, 2:1, etc.)
Blinding

- **Blinding** = treatment masking
- Treatment masked from the patient - single blind
- Treatment masked from both the patient and the study personnel - double blind
- Blinding controls for
  - Treatment/procedure bias
  - Assessment bias
- No blinding for members of DSMB
- Blinding isn’t always possible
  - Drugs being compared have different modes of delivery - infusion versus tablet
  - Blinding in trials of devices or surgical procedures is difficult or impossible
Study populations

*Intention-to-treat* (ITT) is the idea that patients in a randomized clinical trial should be analyzed as part of the treatment group to which they were assigned, even if they did not actually receive the intended treatment. For assessing efficacy, analysis of the ITT population is preferred.

*Treatment received* (TR) is the idea that patients should be analyzed according to the treatment actually given, even if the randomization calls for something else. For assessing safety, analysis of the TR population is preferred. This is sensible since we want to attribute any severe adverse events to the treatment actually received.
Efficacy based on the ITT population

- Many factors contribute to a patient’s inability to complete the intended therapy or patient adherence
  - Side effects
  - Disease progression
  - Patient/physician preference for a different treatment

- Failure to complete therapy as randomized is almost always an outcome of the study itself

- If ITT population is not analyzed, can result in post-entry selection bias

- Results from efficacy analysis using ITT population is a test of treatment policy/program effectiveness
Correlative biomarker studies

- Secondary or exploratory aims of trial
- Examine association of predictive marker with clinical endpoint
- Example (used with permission) from M Regan, DFCI
- Phase II trial to evaluate response of patients with muscle-invasive urothelial cancer treated with neoadjuvant dose-dense methotrexate, vinblastine, doxorubicin, cisplatin (ddMVAC), followed by radical surgery with curative intent (Choueiri, ASCO 2013, 4530)
- Single-arm, 2-stage design (futility look) - $\alpha = 0.1$, power = 0.85, $n = 37$
- Correlative objective: Investigate tumor expression levels of DNA repair genes (e.g. ERCC1) in relation to response
- High tumor tissue levels of ERCC1 mRNA have been associated with clinical resistance to cisplatin-based chemotherapy in ovarian, gastric, cervical, colon and NSCLC patients
ddMVAC example: ideal versus reality

Result the investigator anticipates: large, clear difference
ddMVAC example: ideal versus reality

Ideal (n=37) 55% response

Less ideal (n=37) 55% response

Reality (n=27) successful assays 55% response

P<.001 P=.03 P=.24

Statistician’s anticipation: smaller difference, greater variability
ddMVAC example: ideal versus reality

 Ideal (n=37)  
55% response

Reality (n=37)  
55% response

Reality (n=27) successful assays  
55% response