



MRC  
Biostatistics  
Unit



UNIVERSITY OF  
CAMBRIDGE

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# **Adaptive Methods in Clinical Research**

## *Lecture 2: Group Sequential Designs*

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1. Introduction to group sequential designs
2. Type I error rate, power and expected sample size
3. Stopping boundaries

# Planning an RCT

- A randomised controlled trial (RCT) is carried out to compare the effectiveness of a new experimental treatment versus a control (placebo or standard treatment).
- Generally, frequentist operating characteristics are controlled: the type I error rate  $\alpha$ , and power  $(1 - \beta)$
- Let  $\theta$  denote some measure of the difference between the effectiveness of the new treatment and the control.
- Testing the null hypothesis  $H_0 : \theta \leq 0$  against the alternative  $H_1 : \theta > 0$ , using a suitable test statistic  $Z$
- *Fixed sample test*: reject  $H_0$  if  $Z > c$ , where  $P(Z > c) \leq \alpha$  under  $H_0$ . Choose sample size  $n$  so that  $P(Z > c) \geq 1 - \beta$  when  $\theta = \delta$

# Group sequential designs

- Consider a group sequential design with a total of  $J$  analyses.
- At  $j$ th analysis, test statistic  $Z_j$  is calculated using patients assessed so far.
- A general one-sided group sequential test is defined by constants  $(l_j, u_j)$  with  $l_j < u_j$  for  $j = 1, \dots, J$  and  $l_J = u_J$

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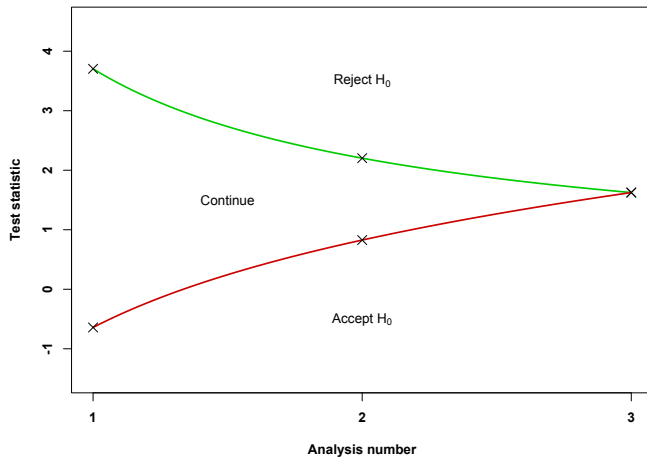
After group  $j = 1, \dots, J - 1$

- |                   |   |
|-------------------|---|
| if $Z_j \geq u_j$ | stop, reject $H_0$ ( <i>early stopping for efficacy</i> )               |
| if $Z_j \leq l_j$ | stop, do not reject $H_0$ ( <i>early stopping for lack of benefit</i> ) |
| otherwise         | continue to group $j + 1$   |

after group  $J$

- |                   |                           |
|-------------------|---------------------------|
| if $Z_J \geq u_J$ | stop, reject $H_0$        |
| if $Z_J < l_J$    | stop, do not reject $H_0$ |

# Group sequential design schematic



# Repeated testing of $H_0$

- A naive way to apply group sequential designs: test  $H_0$  at significance level  $\alpha$  several times throughout the trial.
- E.g.,  $u_1 = \dots = u_J = 1.64$ ,  $l_1 = \dots$ ,  $l_{J-1} = -\infty$ ,  $l_J = 1.64$
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Number of analyses	Type I error rate
1	0.050
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- This inflates the probability of making a type I error:

Number of analyses	Type I error rate
1	0.050
2	0.080
3	0.101
5	0.130
10	0.172

- Need to adjust the critical value used to ensure the overall type I error rate is controlled.

- A long history: theory dates back to Wald in the 1940s, with early medical applications by Armitage in the 1950s/60s
- Now one of the most commonly used type of adaptive designs

# Calculating error probabilities

- Let  $\mathbf{l} = (l_1, \dots, l_J)$  be the *lack of benefit boundaries* and  $\mathbf{u} = (u_1, \dots, u_J)$  the *efficacy boundaries*.
- Let  $n_j$  denote the number of patients who have been assessed on each arm by the  $j$ th analysis, and let  $\mathbf{n} = (n_1, \dots, n_J)$

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- Let  $n_j$  denote the number of patients who have been assessed on each arm by the  $j$ th analysis, and let  $\mathbf{n} = (n_1, \dots, n_J)$
- Problem: how to choose  $\mathbf{l}$ ,  $\mathbf{u}$  and  $\mathbf{n}$  so that design has overall type I error rate  $\alpha$  and power  $(1 - \beta)$
- To solve this we can solve a simpler problem: for a trial design with parameters  $\mathbf{l}$ ,  $\mathbf{u}$  and  $\mathbf{n}$ , what is the type I error rate and power?
- A useful theorem can be used.

# Calculating error probabilities

- Let  $\hat{\theta}_j$  be the maximum likelihood estimate (MLE) of  $\theta$  at analysis  $j$
- Let  $\mathcal{I}_j$  denote the (Fisher) *information* at analysis  $j$ 
  - ▶ Asymptotic variance of MLE given by  $\text{var}(\hat{\theta}_j) = \frac{1}{\mathcal{I}_j}$
- At analysis  $j$ , the Wald statistic,  $Z_j$  is calculated:

$$Z_j = \frac{\hat{\theta}_j}{\sqrt{\text{var}(\hat{\theta}_j)}} = \hat{\theta}_j \sqrt{\mathcal{I}_j} \quad (1)$$

# Calculating error probabilities

- **Theorem 1:** the asymptotic joint distribution of  $(Z_1, \dots, Z_J)$  given  $(\mathcal{I}_1, \dots, \mathcal{I}_J)$  has the following properties:
  - ▶  $(Z_1, \dots, Z_J)$  is multivariate normal
  - ▶  $E(Z_j) = \theta \sqrt{\mathcal{I}_j}$
  - ▶  $\text{Cov}(Z_{j_1}, Z_{j_2}) = \sqrt{\mathcal{I}_{j_1} \mathcal{I}_{j_2}}$  for  $1 \leq j_1 \leq j_2 \leq J$

For proof (and required regularity conditions), see Jennison and Turnbull (JASA 1997)

- **Key message:** Group sequential design theory is applicable in a very wide variety of design scenarios
  - ▶ E.g. Binary outcomes, time-to-event outcomes, GLMs, ...

## Example: normally distributed endpoint

- Responses  $Y_{0i} \sim N(\mu_0, \sigma^2)$  for patients on control treatment, and  $Y_{1i} \sim N(\mu_1, \sigma^2)$  for patients on experimental treatment. Assume  $\sigma^2$  known.
- Parameter  $\theta = \mu_1 - \mu_0$  is of interest.

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- Parameter  $\theta = \mu_1 - \mu_0$  is of interest.
- At analysis  $j$ , MLE  $\hat{\theta}_j$  is

$$\bar{Y}_1^{(j)} - \bar{Y}_0^{(j)} = \left( \frac{1}{n_j} \sum_{i=1}^{n_j} Y_{1i} - \frac{1}{n_j} \sum_{i=1}^{n_j} Y_{0i} \right) \sim N\left(\theta, \frac{2\sigma^2}{n_j}\right)$$

- Wald test statistic is  $Z_j = (\bar{Y}_1^{(j)} - \bar{Y}_0^{(j)})\sqrt{\mathcal{I}_j}$ , where  $\mathcal{I}_j = \frac{n_j}{2\sigma^2}$
- $(Z_1, \dots, Z_J)$  is multivariate normal (since linear combination of independent normals), and marginally  $Z_j \sim N(\theta\sqrt{\mathcal{I}_j}, 1)$



## Example: normally distributed endpoint

- $(Z_1, \dots, Z_J)$  is multivariate normal (since linear combination of independent normals), and  $Z_j \sim N(\theta\sqrt{\mathcal{I}_j}, 1)$
- For  $j_1 \leq j_2$ ,

$$\begin{aligned}\text{Cov}(Z_{j_1}, Z_{j_2}) &= \text{Cov}\left(\bar{Y}_1^{(j_1)} - \bar{Y}_0^{(j_1)}, \bar{Y}_1^{(j_2)} - \bar{Y}_0^{(j_2)}\right) \sqrt{\mathcal{I}_{j_1}} \sqrt{\mathcal{I}_{j_2}} \\ &= \frac{2}{n_{j_1} n_{j_2}} n_{j_1} \sigma^2 \sqrt{\mathcal{I}_{j_1}} \sqrt{\mathcal{I}_{j_2}} = \sqrt{\mathcal{I}_{j_1} / \mathcal{I}_{j_2}}\end{aligned}$$

## Example: normally distributed endpoint

- Using the multivariate normal distribution of  $(Z_1, \dots, Z_J)$ , we can derive the probability of the trial stopping at each stage.

## Example: normally distributed endpoint

- Using the multivariate normal distribution of  $(Z_1, \dots, Z_J)$ , we can derive the probability of the trial stopping at each stage.
- For example, the probability of stopping for efficacy in the second stage is:

$$\int_{l_1}^{u_1} \int_{l_2}^{\infty} \phi_2((y_1, y_2), \left( \theta \sqrt{\frac{n_1}{2\sigma^2}}, \theta \sqrt{\frac{n_2}{2\sigma^2}} \right), \begin{pmatrix} 1 & \sqrt{n_1/n_2} \\ \sqrt{n_1/n_2} & 1 \end{pmatrix}) dy_2 dy_1$$

where  $\phi_2(y, \mu, \Sigma)$  is the density of the bivariate normal distribution with mean  $\mu$  and covariance  $\Sigma$  at  $y$ .

## Example: normally distributed endpoint

- To get type I error rate, add up probabilities of stopping for efficacy at each analysis when  $\theta = 0$
- To get power, add up probabilities of stopping for efficacy at each analysis when  $\theta = \delta$
- E.g. for three-stage design with  $\mathbf{u} = (2.5, 2, 1.5)$ ,  $\mathbf{l} = (0, 0.75, 1.5)$ ,  $\mathbf{n} = (20, 40, 60)$ ,  $\delta = 0.5$ ,  $\sigma^2 = 1$ :

Analysis	$\theta = 0$		$\theta = 0.5$	
	Prob for futility	Prob stop for efficacy	Prob for futility	Prob stop for efficacy
1	0.500	0.006	0.057	0.179
2	0.299	0.019	0.042	0.420
3	0.137	0.038	0.049	0.253

So type I error rate =  $0.006 + 0.019 + 0.038 = 0.063$ ;

power =  $0.179 + 0.42 + 0.253 = 0.852$ .

# Finding a design with given error rates

- We want to find a design with given type I error rate and power.
- Generally it is assumed that analyses are equally spaced in terms of patients, so that  $(n_1, \dots, n_J) = (n, 2n, \dots, Jn)$ 
  - ▶ In this case, type I error rate is independent of  $n$
- First find boundary that gives correct type I error rate, then find sample size  $n$  to give correct power.

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- First find boundary that gives correct type I error rate, then find sample size  $n$  to give correct power.
- One approach: choose  $\mathbf{u}$  and  $\mathbf{l}$ , then find  $c$  such that  $(cl_1, \dots, cl_J)$  and  $(cu_1, \dots, cu_J)$  gives correct type I error rate.

## Example:

- Assume normally distributed endpoint as before
- For  $\mathbf{l} = (0, 0.75, 1.5)$  and  $\mathbf{u} = (2.5, 2, 1.5)$ ,  $c = 1.081$  gives type I error rate equal to 0.05

## Example (continued):

- Stopping boundaries  $\mathbf{l} = (0, 0.811, 1.622)$  and  $\mathbf{u} = (2.703, 2.162, 1.622)$
- In this case,  $n = 26$  is needed for 90% power when  $\theta = 0.5$ .
- So  $\mathbf{n} = (26, 52, 78)$ ,  $\mathbf{l} = (0, 0.811, 1.622)$  and  $\mathbf{u} = (2.703, 2.162, 1.622)$  is a group sequential design with type I error rate 0.05 and power 0.9.

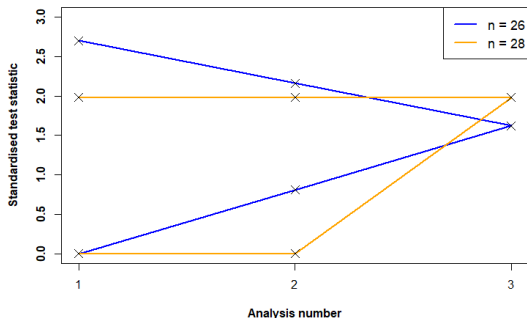
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- However,  
 $\mathbf{n} = (28, 56, 84)$ ,  $\mathbf{l} = (0, 0, 1.98)$ ,  $\mathbf{u} = (1.98, 1.98, 1.98)$  is another design with the same characteristics



# Finding a design with given error rates

## Example (continued):



- In fact there are an *infinite* number of possible designs
- How to pick between them?

## Expected sample size

- One way to distinguish between designs with the same type I error rate and power is through *expected sample size* (ESS).

## Expected sample size

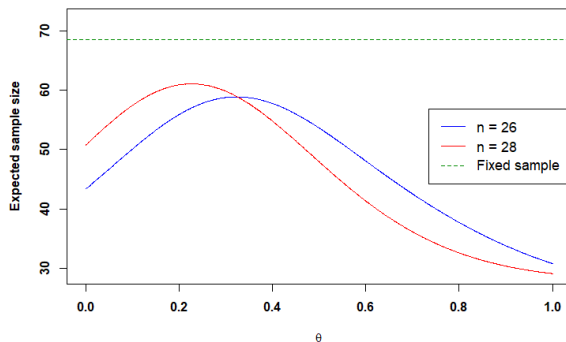
- One way to distinguish between designs with the same type I error rate and power is through *expected sample size* (ESS).
- Can be calculated using the stopping probabilities. For example, from previous example when  $\mathbf{n} = (20, 40, 60)$  and  $\theta = 0.5$ :

Analysis	Prob stop for futility	Prob stop for efficacy	Total prob of stopping
1	0.500	0.006	0.506
2	0.299	0.019	0.318
3	0.137	0.038	0.176

- The ESS is:  $(0.506 \times 20) + (0.318 \times 40) + (0.176 \times 60) = 33.4$
- More generally, the ESS is  $\sum_{j=1}^J n_j p_j$ , where  $p_j$  is the probability of stopping at analysis  $j$

# Expected sample size

For example, expected sample size for the two designs mentioned on slide 16.



Which is preferable?

## Expected and maximum sample size

- If trial is powered to detect difference  $\theta = 0.5$ , the true  $\theta$  may be smaller than this.
- Thus design in blue may be preferable in practice – however, depends on prior beliefs on  $\theta$
- Another advantage of the blue design is that its *maximum sample size*, MSS, is lower (78 vs 84).
  - ▶ This is the number of patients recruited if the trial does not stop until the last analysis
  - ▶ For fixed sample size trial, need 69 per arm.

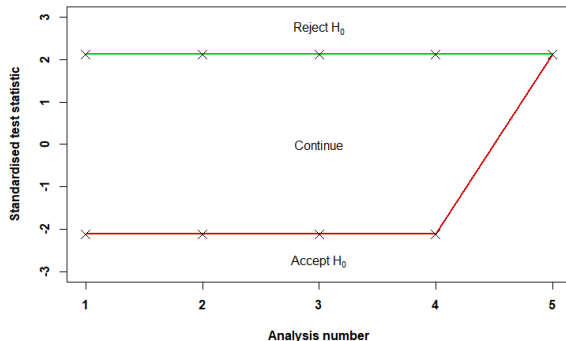
# Choosing stopping boundary shapes

- The shape of the stopping boundaries will determine the ESS and MSS.
- Two main ways to choose between the infinite number of possible shapes:
  1. Use some 'fixed' boundary shape (specified through a simple functional form);
  2. Search for an optimal design to (e.g.) minimise ESS for some value of  $\theta$ .
- First method is much quicker, but second method allows greater control over the ESS properties of the design.
- Common boundary shapes to choose from:
  1. Pocock
  2. O'Brien-Fleming
  3. Triangular test

# Common boundary shapes

## Pocock

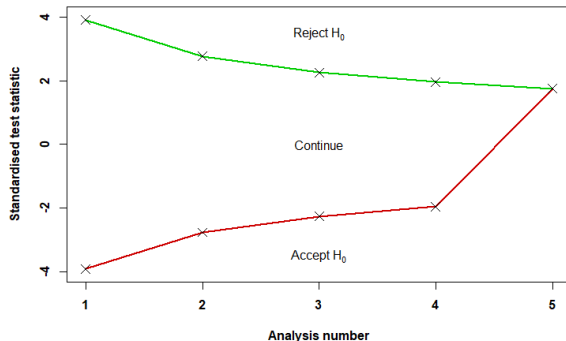
- $u_1 = \dots = u_J = C$
- $l_1 = \dots = l_{J-1} = -C, l_J = C$
- $C$  chosen to ensure correct type I error rate and power



# Common boundary shapes

## O'Brien-Fleming

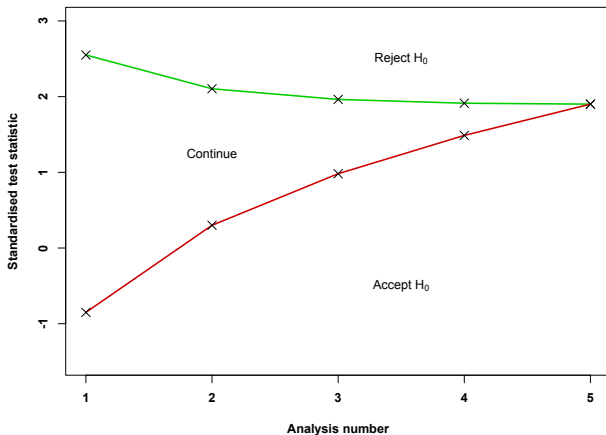
- $u_j = C\sqrt{J/j}, j = 1, \dots, J$
- $l_j = -C\sqrt{J/j}, j = 1, \dots, J-1$  and  $l_J = C$
- $C$  chosen to ensure correct type I error rate and power





# Choosing stopping boundary shapes

## Triangular test



# Optimal designs

- An alternative way of choosing stopping boundary shapes is *optimal designs*.
- Firstly, define a *feasible design* as a group sequential design which meets some type I error rate and power constraints.
- An optimal design is feasible *and* minimises the expected sample size for some value of  $\theta$ .
- Infinite number of optimal designs; some commonly discussed ones:
  - ▶ Null-optimal design; optimal for  $\theta = 0$ ;
  - ▶ Alternative-optimal design; optimal for  $\theta = \delta$
  - ▶  $\theta$ -minimax design; has the lowest maximum expected sample size.

## (Near-)Optimal boundaries

- To aid finding optimal design, can consider a constrained set of possible boundary shapes:
- Stopping boundaries are

$$u_j = C_u(j/J)^{\Delta_u-1/2}$$
$$l_j = \delta\sqrt{\mathcal{I}_j} - C_l(j/J)^{\Delta_l-1/2}$$

- The constants  $C_u$  and  $C_l$  are chosen to ensure correct type I error and power
- Can then optimise  $(\Delta_u, \Delta_l)$
- This is the approach taken in the `OptGS` package (see Practical)

# Advantages and disadvantages of group sequential trials

- Advantages:
  - ▶ Fewer patients required on average compared to fixed sample-size designs.
  - ▶ If one treatment is ineffective, fewer patients on average will be exposed to it.
  - ▶ Reduces time to get effective treatment to market.
- Disadvantages:
  - ▶ If trial continues to the end, more patients will be used compared to fixed sample-size design.
  - ▶ Interim analyses can introduce practical issues
  - ▶ Complicates the analysis (see Lecture 6 ...)

1. Jennison, C. and Turnbull, B. (2000). Group Sequential Methods with Applications to Clinical Trials. Chapman & Hall/CRC.
2. Wason, J. Mander, A. and Thompson, S. (2012). Optimal multi-stage designs for randomised clinical trials with continuous outcomes. Statistics in Medicine, 31, 301-312.