Note: The corresponding sample size procedure, found in PASS Sample Size software, is *Group-Sequential Superiority by a Margin Tests for Two Hazard Rates (Simulation)*.

# Introduction

This procedure is used to test superiority by a margin for the difference of two hazard rates in stages (sometimes called looks or interim analyses) using group-sequential methods. This methodology assumes an underlying Exponential model. Unless the stage boundaries are entered directly, the stage boundaries are defined using a specified spending function. Futility boundaries can be binding or non-binding. Futility boundaries are specified through a beta-spending function.

Sample size re-estimation, based on current-stage sample sizes and parameter estimates, may also be obtained in this procedure.

The spending functions available in this procedure are the O'Brien-Fleming analog, the Pocock analog, the Hwang-Shih-DeCani gamma family, and the power family.

At each stage, the current and future boundaries are calculated based on the accumulated information proportion. Conditional and predictive power for future stages is also given.

A group-sequential boundary and analysis graph is produced in this procedure.



At each stage, stage-adjusted difference estimates, confidence intervals, and p-values are available.

The probabilities of crossing future boundaries may also be assessed, using simulation.

The format of the data for use in this procedure is three columns: one column for the response values, one column defining the two groups, and a third column defining the stage.

# **Outline of a Group-Sequential Study**

There are three basic phases of a group-sequential (interim analysis) study:

- Design
- Group-Sequential Analysis
- Reporting

### **Design Phase – Determine the Number of Subjects**

To begin the group-sequential testing process, an initial calculation should be made to determine the sample size and target information if the final stage is reached (maximum information). The sample size calculation requires the specification of the following:

- Alpha
- Power
- Test Direction
- Types of boundaries (efficacy, binding futility, non-binding futility)
- Maximum number of stages
- Proportion of maximum information at each stage
- Spending functions
- Assumed survival rates

The design phase calculation may be done in the **PASS** sample size software program. **PASS** software permits the user to easily try a range of hazard rate differences, as these values are typically not known in advance.

The resulting sample size of the sample size calculation also permits the calculation of the maximum information, which is the total information of the study if the final stage is reached (for calculation details, see the Information section later in this chapter).

Based on the maximum information, the target information and target sample size of each stage may be calculated. In particular, this permits the user to have a target sample size for the first stage.

Although it is likely to change over the course of the group-sequential analysis, a design group-sequential boundary plot can be a useful visual representation of the design:



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### **Group-Sequential Analysis Phase**

A group sequential analysis consists of a series of stages where a decision to stop or continue is made at each stage.

### First Interim Stage

The design phase gives the target number of subjects for the first stage, based on the time of the first stage and the accrual specification. The study begins, and response data is collected for subjects, moving toward the first-stage target number of subjects, until a decision to perform an analysis on the existing data is made. The analysis at this point is called the first stage.

Unless the number of subjects at the first stage matches the design target for the first stage, the calculated information at the first stage will not exactly match the design information for the first stage. Generally, the calculated information will not differ too greatly from the design information, but regardless, spending function group-sequential analysis is well-suited to make appropriate adjustments for any differences.

The first stage information is divided by the maximum information to obtain the stage one information proportion (or information fraction). This information proportion is used in conjunction with the spending function(s) to determine the alpha and/or beta spent at that stage. In turn, stage one boundaries, corresponding to the information proportion, are calculated.

A *z*-statistic is calculated from the raw hazard rate difference, or other logrank-type method. The stage one *z*-statistic is compared to each of the stage one boundaries. Typically, if one of the boundaries is crossed, the study is stopped (non-binding futility boundaries may be an exception).



If none of the boundaries are crossed the study continues to the next stage.



If none of the boundaries are crossed it may also be useful to examine the conditional power or stopping probabilities of future stages. Conditional power and stopping probabilities are based on the user-specified supposed true difference.

### Second and other interim stages (if reached)

If the first stage time proportion is not equal to the design time proportion, a designation must be made at this point as to the target time of the second stage. Two options are available in this procedure.

One option is to target the information proportion of the original design. For example, if the original design proportions of a four-stage design are 0.25, 0.50, 0.75, 1.0, and the stage one observed proportion is 0.22, the researcher might still opt to target 0.50 for the second stage, even though that now requires an additional information accumulation of 0.28 (proportion). The third and fourth stage targets would also remain 0.75 and 1.0.

A second option is to adjust the target information proportionally to the remaining proportions. For this option, if the design proportions are 0.25, 0.50, 0.75, 1.0, and 0.22 is observed, the remaining 0.78 is distributed proportionally to the remaining stages. In this example, the remaining target proportions become 0.48, 0.74, 1.0.

For either option, once the target information is determined for the next stage, revised target sample sizes are given, and the study continues until the decision is made to perform the next interim analysis on the cumulative response data. In the same manner as the first stage, the current stage information proportion is used with the spending function to determine alpha and/or beta spent at the current stage. The current stage boundaries are then computed. The *z*-statistic is calculated and compared to the boundaries, and a decision is made to stop or continue.

If a boundary is crossed, the study is typically stopped.



If none of the boundaries are crossed the study continues to the next stage.



Once again, if no boundary is crossed, conditional power and stopping probabilities may be considered based on a choice of a supposed true difference.

The study continues from stage to stage until the study is stopped for the crossing of a boundary, or until the final stage is reached.

### Final Stage (if reached)

The final stage (if reached) is similar to all the interim stages, with a couple of exceptions. For all interim analyses the decision is made whether to stop for the crossing of a boundary, or to continue to the next stage. At the final stage, only the decision of efficacy or futility can be made.

Another intricacy of the final stage that does not apply to the interim stages is the calculation of the maximum information. At the final stage, the current information must become the maximum information, since the spending functions require that the proportion of information at the final look must be 1.0. If the current information at the final stage is less than the design maximum information, the scenario is sometimes described as *under-running*. Similarly, if the current information at the final stage is greater than the design maximum information, the result may be termed *over-running*.

For both under-running and over-running, the mechanism for adjustment is the same, and is described in the Technical Details section, under Information and Total Information.

Aside from these two exceptions, the final stage analysis is made in the same way that interim analyses were made. The remaining alpha and beta to be spent are used to calculate the final stage boundaries. If the test is a one-sided test, then the final stage boundary is a single value. The final stage *z*-statistic is computed from the sample hazard rates of the complete data from each group. The *z*-statistic is compared to the boundary and a decision of efficacy or futility is made.

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# **Reporting Phase**

Once a group-sequential boundary is crossed and the decision is made to stop, there remains the need to properly summarize and communicate the study results. Some or all of the following may be reported:

- Boundary plot showing the crossed boundary
- Adjusted confidence interval and estimate of the hazard rate difference
- Sample size used

### Boundary plot showing the crossed boundary

The boundary plot gives an appropriate visual summary of the process leading to the reported decision of the study.



### Adjusted confidence interval and estimate of the hazard rate difference

Due to the bias that is introduced in the group-sequential analysis process, the raw data confidence interval of the difference in hazard rates should not be used. An adjusted confidence interval should be used instead. See the Adjusted Confidence Interval topic of the Technical Details section for details. The mid-point of the adjusted confidence interval may be useful as a point estimate of the difference in hazard rates. Further, the confidence level at which the adjusted confidence interval limit is zero gives a rough adjusted *p*-value.

### Sample size used

The sample size at the point the study was stopped should be reported in addition to the sample size that would have been used had the final stage been reached.

# **Technical Details**

Many articles and texts have been written about group sequential analysis. Details of many of the relevant topics are discussed below, but this is not intended to be a comprehensive review of group-sequential methods. One of the more influential works in the area of group-sequential analysis is Jennison and Turnbull (2000).

## **Null and Alternative Hypotheses**

For superiority by a margin tests of two hazard rates, the appropriate null and alternative hypotheses depend on whether higher hazard rates are better or higher hazard rates are worse.

### Case 1: Low Hazard Rates Good

In this case, lower hazard rates are better. The hypotheses are arranged so that rejecting the null hypothesis implies that the treatment hazard rate is less than the reference hazard rate by at least the margin of superiority. The value of  $\delta$  at which power is calculated must be less than  $-|M_S|$ . The null and alternative hypotheses with  $\delta_0 = -|M_S|$  are

$H_0: h_1(T) \ge h_2(T) -  M_S $	versus	$H_1: h_1(T) < h_2(T) -  M_S $
$H_0: h_1(T) - h_2(T) \ge - M_S $	versus	$H_1: h_1(T) - h_2(T) < - M_S $
$H_0: \delta \ge - M_S $	versus	$H_1:\delta<- M_S $

### Case 2: High Hazard Rates Good

In this case, higher hazard rates are better. The hypotheses are arranged so that rejecting the null hypothesis implies that the treatment hazard rate is greater than the reference hazard rate by at least the margin of superiority. The value of  $\delta$  at which power is calculated must be greater than  $|M_S|$ . The null and alternative hypotheses with  $\delta_0 = |M_S|$  are

$H_0: h_1(T) \le h_2(T) +  M_S $	versus	$H_1: h_1(T) > h_2(T) +  M_S $
$H_0: h_1(T) - h_2(T) \le  M_S $	versus	$H_1: h_1(T) - h_2(T) >  M_S $
$H_0:\delta \le  M_S $	versus	$H_1: \delta >  M_S $

## **Stages in Group-Sequential Testing**

The potential to obtain the benefit from a group-sequential design and analysis occurs when the response data are collected over a period of weeks, months, or years rather than all at once. A typical example is the case where patients are enrolled in a study as they become available, as in many types of clinical trials.

A group-sequential testing stage is a point in the accumulation of the data where an interim analysis occurs, either by design or by necessity. At each stage, a test statistic is computed with all the accumulated data, and it is determined whether a boundary (efficacy or futility) is crossed. When an efficacy (or futility) boundary is crossed, the study is usually concluded, and inference is made. If the final stage is reached, the group-sequential design forces a decision of efficacy or futility at this stage.

For the discussions below, a non-specific interim analysis stage is referenced as *k*, and the final stage is *K*.

### Test Statistic (MLE)

The *z*-statistic from MLE estimates for any stage *k* is obtained from all the accumulated data up to and including that stage.

The general form of the test statistic is, when lower hazard rates are better:

$$z_{k} = \frac{\hat{h}_{1k} - \hat{h}_{2k} - (-|SM|)}{\sqrt{\hat{\sigma}_{k}^{2}(\hat{h}_{1k}) + \hat{\sigma}_{k}^{2}(\hat{h}_{2k})}} = \frac{\hat{h}_{1k} - \hat{h}_{2k} + |SM|}{\sqrt{\hat{\sigma}_{k}^{2}(\hat{h}_{1k}) + \hat{\sigma}_{k}^{2}(\hat{h}_{2k})}}$$

and, when higher hazard rates are better:

$$z_{k} = \frac{\hat{h}_{1k} - \hat{h}_{2k} - |SM|}{\sqrt{\hat{\sigma}_{k}^{2}(\hat{h}_{1k}) + \hat{\sigma}_{k}^{2}(\hat{h}_{2k})}}$$

with

$$\hat{h}_{ik} = \frac{\sum_{j=1}^{n_{ik}} c_{ijk}}{\sum_{j=1}^{n_{ik}} x_{ijk}}$$

$$\hat{\sigma}_{k}^{2}(\hat{h}_{ik}) = \frac{\sum_{j=1}^{n_{ik}} c_{ijk}}{\left(\sum_{j=1}^{n_{ik}} x_{ijk}\right)^{2}} = \frac{\hat{h}_{ik}^{2}}{\sum_{j=1}^{n_{ik}} c_{ijk}}$$

where

i = 1,2 for the two groups

 $\hat{h}_{ik}$  is the estimated group hazard rate at stage k

 $\hat{\sigma}_k^2(\hat{h}_{ik})$  is the variance of the hazard rate estimator

 $c_{ijk}$  is an indicator of censoring

 $x_{ijk}$  is the elapsed time

### Group-Sequential Design Phase

In most group-sequential studies there is a design or planning phase prior to beginning response collection. In this phase, researchers specify the anticipated number and spacing of stages, the types of boundaries that will be used, the desired alpha and power levels, the spending functions, and the anticipated hazard rates with the corresponding estimate of the true difference in hazard rates.

Based on these input parameters, an initial set of boundaries is produced, an estimate of the total number of needed subjects is determined, and the anticipated total information at the final stage is calculated. The appropriate procedure in **PASS** (sample size software) can be used to make these planning phase sample size estimation calculations.

### Information and Total Information

In the group-sequential design phase, the final stage (*K*) or total (design) information is calculated from the specified rates and the final sample sizes, as

$$I_K^* = \frac{1}{\frac{\sigma_K^2(h_1, l_1, p_1)}{n_{1K}} + \frac{\sigma_K^2(h_2, l_2, p_2)}{n_{2K}}}$$

where

i = 1,2 for the two groups

 $\sigma_K^2(h_i, l_i, p_i)$  is the variance of the hazard rate estimator

 $h_i$  is the group hazard rate

 $l_i$  is the group loss hazard rate

 $p_i$  is the group patient entry parameter

and

$$\sigma_K^2(h_i, l_i, p_i) = h_i^2 \left( \frac{h_i}{h_i + l_i} + \frac{h_i p_i e^{-(h_i + l_i)T} \left( 1 - e^{(h_i + l_i - p_i)T_0} \right)}{(1 - e^{-p_i T_0})(h_i + l_i)(h_i + l_i - p_i)} \right)^{-1}$$

for *i* = 1,2,

where

 $T_0$  is the accrual time

*T* is the total time

If patient entry is uniform, the group variance is (Lachin and Foulkes, 1986):

$$\sigma_K^2(h_i, l_i, p_i) = h_i^2 \left( \frac{h_i}{h_i + l_i} \left[ 1 - \frac{e^{-(T - T_0)(h_i + l_i)} - e^{-T(h_i + l_i)}}{T_0(h_i + l_i)} \right] \right)^{-1}$$

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The information at any data stage *k* may be calculated from the specified rates and the sample sizes, as

$$I_k = \frac{1}{\hat{\sigma}_k^2(\hat{h}_{1k}) + \hat{\sigma}_k^2(\hat{h}_{2k})}$$

with variance estimates as defined in the Test Statistic (MLE) section.

The proportion of the total information (or information fraction) at any stage is

$$p_k = \frac{I_k}{I_K^*}$$

The information fractions are used in conjunction with the spending function(s) to define the alpha and/or beta to be spent at each stage.

To properly use the spending function at the final stage, it is required that  $p_K = 1$ . However, if the final stage is reached, we see that

$$I_{K} = \frac{1}{\hat{\sigma}_{K}^{2}(\hat{h}_{1K}) + \hat{\sigma}_{K}^{2}(\hat{h}_{2K})} \neq I_{K}^{*} = \frac{1}{\frac{\sigma_{K}^{2}(h_{1}, l_{1}, p_{1})}{n_{1K}} + \frac{\sigma_{K}^{2}(h_{2}, l_{2}, p_{2})}{n_{2K}}}$$

so that

$$p_K = \frac{I_K}{I_K^*} \neq 1$$

When  $I_K > I_K^*$ , it is called over-running. When  $I_K < I_K^*$ , it is called under-running. In either case, the spending function is adjusted to accommodate the inequality, by redefining

 $I_K^* = I_K$ 

See the discussion in Wassmer and Brannath (2016), pages 78-79, or Jennison and Turnbull (2000), pages 153-154, 162.

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### **Types of Boundaries**

A variety of boundary designs are available to reflect the needs of the study design.

### **Efficacy Only**

The simplest group-sequential test involves a single set of stage boundaries with early stopping for efficacy.



### **Efficacy and Binding Futility**

This design allows early stopping for either efficacy or futility. For binding futility designs, the Type I error protection (alpha) is only maintained if the study is strictly required to stop if either boundary is crossed.



### **Efficacy and Non-Binding Futility**

This design also allows early stopping for either efficacy or futility. For non-binding futility designs, the Type I error protection (alpha) is maintained, regardless of whether the study continues after crossing a futility boundary. However, the effect is to make the test conservative (alpha is lower than the stated alpha and power is lower than the stated power).



### **Futility Only**

In this design, the interim analyses are used only for futility. Please be aware that, due to computational complexity, these boundaries may take several minutes to compute, particularly when some stages are skipped.



### **Boundary Calculations**

The foundation of the spending function approach used in this procedure is given in Lan & DeMets (1983). This procedure implements the methods given in Reboussin, DeMets, Kim, & Lan (1992) to calculate the boundaries and stopping probabilities of the various group sequential designs. Some adjustments are made to these methods to facilitate the calculation of futility boundaries.

## **Binding vs. Non-Binding Futility Boundaries**

Futility boundaries are used to facilitate the early stopping of studies when early evidence leans to lack of efficacy. When binding futility boundaries are to be used, the calculation of the futility and efficacy boundaries assumes that the study will be strictly stopped at any stage where a futility or efficacy boundary is crossed. If strict adherence is not maintained, then the Type I and Type II error probabilities associated with the boundaries are no longer valid. One (perhaps undesirable) effect of using binding futility boundaries is that the resulting final stage boundary may be lower than the boundary given in the corresponding fixed-sample design.

When non-binding futility boundaries are calculated, the efficacy boundaries are first calculated ignoring futility boundaries completely. This is done so that alpha may be maintained whether or not a study continues after crossing a futility boundary. One (perhaps undesirable) effect of using non-binding futility boundaries is that the overall group-sequential test becomes conservative (alpha is lower than the stated alpha and power is lower than the stated power).

### **Spending Functions**

Spending functions are used to distribute portions of alpha (or beta) to the stages according to the proportion of accumulated information at each look.

### **Spending Function Characteristics**

• Spending functions give a value of zero when the proportion of accumulated information is zero.

 $\alpha(0) = 0$  (for alpha-spending)

 $\beta(0) = 0$  (for beta-spending)

- Spending functions are increasing functions.
- Spending functions give a value of alpha (or beta) when the proportion of accumulated information is one.

 $\alpha(1) = \alpha$  (for alpha-spending)  $\beta(1) = \beta$  (for beta-spending)

Using spending functions in group-sequential analyses is very flexible in that neither the information proportions nor the number of stages need be specified in advance to maintain Type I and Type II error protection.

### Spending Functions Available in this Procedure

The following spending functions are shown as alpha-spending functions. The corresponding beta-spending function is given by replacing  $\alpha$  with  $\beta$ .

### **O'Brien-Fleming Analog**

The O'Brien Fleming Analog (Lan & DeMets, 1983) roughly mimics the O'Brien-Fleming (non-spending function) design, with the key attribute that only a small proportion of alpha is spent early. Its popularity comes from it proportioning enough alpha to the final stage that the final stage boundary is not too different from the fixed-sample (non-group-sequential) boundary.

$$\alpha(0) = 0$$
  

$$\alpha(p_k) = 2 - 2\Phi\left(\frac{Z_{1-\alpha/2}}{\sqrt{p_k}}\right)$$
  

$$\alpha(1) = \alpha$$

![](_page_14_Figure_8.jpeg)

# Pocock Analog

The Pocock Analog (Lan & DeMets, 1983) roughly mimics the Pocock (non-spending function) design, with the key attribute that alpha is spent roughly equally across all stages.

$$\alpha(0) = 0$$
  

$$\alpha(p_k) = \alpha \ln (1 + (e - 1)p_k)$$
  

$$\alpha(1) = \alpha$$

![](_page_15_Figure_5.jpeg)

### **Power Family**

The power family of spending functions has a  $\rho$  parameter that gives flexibility in the spending function shape.

$$\alpha(0) = 0$$
  

$$\alpha(p_k) = p_k^{\rho}, \ \rho > 0$$
  

$$\alpha(1) = \alpha$$

A power family spending function with a  $\rho$  of 1 is similar to a Pocock design, while a power family spending function with a  $\rho$  of 3 is more similar to an O'Brien-Fleming design.

 $\rho = 1$ 

![](_page_16_Figure_4.jpeg)

 $\rho = 2$ 

![](_page_16_Figure_6.jpeg)

 $\rho = 3$ 

![](_page_17_Figure_3.jpeg)

### Hwang-Shih-DeCani (Gamma Family)

The Hwang-Shih-DeCani gamma family of spending function has a  $\gamma$  parameter that allows for a variety of spending functions.

$$\begin{aligned} \alpha(0) &= 0\\ \alpha(p_k) &= \alpha \left( \frac{1 - e^{-\gamma p_k}}{1 - e^{-\gamma}} \right), \ \gamma \neq 0\\ \alpha(p_k) &= \alpha p_k, \ \gamma = 0\\ \alpha(1) &= \alpha \end{aligned}$$

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 $\gamma = -3$ 

![](_page_18_Figure_4.jpeg)

 $\gamma = -1$ 

![](_page_18_Figure_6.jpeg)

 $\gamma = 1$ 

![](_page_19_Figure_3.jpeg)

 $\gamma = 3$ 

![](_page_19_Figure_5.jpeg)

# Adjusted Confidence Intervals

Except at the first stage, the raw (naïve) estimates of the confidence interval limits are inherently biased in the group-sequential analysis setting. The method given in Reboussin, DeMets, Kim, & Lan (1992) is used to calculate appropriately adjusted confidence limits and is based on Kim and DeMets (1987). This stage-wise ordering method is recommended by Jennison and Turnbull (2000) as "the only method available for use with unpredictable information sequences."

The methods used for the calculation of these confidence limits are based on the assumption that the current stage is the stopping stage of the study (typically from the crossing of a boundary).

As the methods in Reboussin *et al.* (1992) give only efficacy boundary results, the calculation adjustments are based only on the efficacy boundaries. The futility boundaries are not used in these calculations, except as they affect the efficacy boundaries.

## Adjusted 'p-values'

As the raw (naïve) *p*-value is inherently biased in the group-sequential analysis setting, a sequential-test adjusted *p*-value should be used instead. Rather than giving a *p*-value in this procedure, a search is used to determine the adjusted confidence interval level at which an interval limit equals zero. As such, the methods used for this calculation are also based on the assumptions that the current stage is the stopping stage of the study. Again, only efficacy bounds are used.

## Adjusted Hazard Rate Difference

The adjusted hazard rate difference is a rough estimate of the difference calculated simply as the midpoint of the adjusted confidence interval limits.

### **Conditional Power**

From Jennison and Turnbull (2000) pages 205 to 208, the general upper one-sided conditional power at stage k for rejecting a null hypothesis about a parameter  $\theta$  at the end of the study, given the observed test statistic,  $Z_k$ , is computed as

$$P_{uk}(\theta) = \Phi\left(\frac{Z_k\sqrt{I_k} - Z_{1-\alpha}\sqrt{I_K} + \theta(I_K - I_k)}{\sqrt{I_K - I_k}}\right),$$

the general lower one-sided conditional power at stage *k* is computed as

$$P_{lk}(\theta) = \Phi\left(\frac{-Z_k\sqrt{I_k} - Z_{1-\alpha}\sqrt{I_K} - \theta(I_K - I_k)}{\sqrt{I_K - I_k}}\right),$$

and the general two-sided conditional power at stage k is computed as

$$P_k(\theta) = \Phi\left(\frac{Z_k\sqrt{I_k} - z_{1-\alpha/2}\sqrt{I_K} + \theta(I_K - I_k)}{\sqrt{I_K - I_k}}\right) + \Phi\left(\frac{-Z_k\sqrt{I_k} - z_{1-\alpha/2}\sqrt{I_K} - \theta(I_K - I_k)}{\sqrt{I_K - I_k}}\right),$$

- $\theta$  = the parameter being tested by the hypothesis
- k = an interim stage at which the conditional power is computed (k = 1, ..., K 1)
- K = the stage at which the study is terminated, and the final test computed
- $Z_k$  = the test statistic calculated from the observed data that has been collected up to stage k
- $I_k$  = the information level at stage k
- $I_K$  = the information level at the end of the study
- $z_{1-\alpha}$  = the standard normal value for the test with a type I error rate of  $\alpha$ .

### **Predictive Power**

*Predictive power* (a Bayesian concept) is the result of averaging the conditional power over the posterior distribution of effect size. From Jennison and Turnbull (2000) pages 210 to 213, the general upper one-sided predictive power at stage *k* is given by

$$P_{uk} = \Phi\left(\frac{Z_k\sqrt{I_K} - Z_{1-\alpha}\sqrt{I_k}}{\sqrt{I_K - I_k}}\right)$$

The general lower one-sided predictive power at stage *k* is given by

$$P_{lk} = \Phi\left(\frac{-Z_k\sqrt{I_K} - Z_{1-\alpha}\sqrt{I_k}}{\sqrt{I_K - I_k}}\right)$$

The general two-sided predictive power at stage *k* is given by

$$P_k = \Phi\left(\frac{|Z_k|\sqrt{I_K} - z_{1-\alpha/2}\sqrt{I_k}}{\sqrt{I_K - I_k}}\right) + \Phi\left(\frac{-|Z_k|\sqrt{I_K} - z_{1-\alpha/2}\sqrt{I_k}}{\sqrt{I_K - I_k}}\right)$$

with all terms defined as in the equations for conditional power.

### Using Simulation to obtain Future Boundary Crossing Probabilities

It can be useful to researchers to know the probability of crossing future group-sequential boundaries, given the data already obtained, and given specified assumed values for the hazard rates. The following steps are used to estimate these probabilities using simulation:

- 1. From the data obtained to the current stage, obtain a current estimate of the information.
- 2. Determine the target (cumulative) sample sizes and numbers of events for each future stage, including the final stage. Fractional sample sizes are rounded up to the next integer.
- 3. For each simulation, append simulated values to the current sample data to obtain a data set with the final stage sample sizes. Simulated values correspond to assumed hazard rates.
- 4. For each data set, determine which boundary or boundaries were crossed first (except in the case of non-binding futility boundaries). The proportion of simulations crossing each boundary (first) provides an estimate of the probability of crossing each boundary, given the specified assumed hazard rates.

### **Non-binding Futility Boundaries**

When non-binding futility boundaries are used, the study may continue when a futility boundary is crossed. The simulation proportions will have a slightly different interpretation when this is the case.

# Data Structure

The data for this procedure is entered in four columns, with an additional column to identify the stage times. Two columns give the individual start and end times. A censor column identifies whether the end time is due to an event or to censoring. Another column identifies the group of the patient, individual, or experimental unit.

These five columns are assigned on the Variables tab of the procedure. Groups 1 and 2 are also assigned on the Variables tab.

The current stage is also identified directly on the Variables tab.

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	1	2	3	4	5	6	7	8	9	
Name	StartDate	EndDate	Censor	Group	Start	End	StageTimes	C8	C9	С
Label										
Data Type	DateTime	DateTime	General	General	General	General	General	General	General	G
Format										
Value Labels										=
Value Order										
Transformation					(StartDate	(EndDate-				
Note										
Filter										-
•										•
	StartDate	EndDate	Censor	Group	<i>⊆f<sub>x</sub></i> Start	<i>⊾f<sub>x</sub></i> End	StageTimes	C8	C9	C
1	1/2/2017	2/12/2017	0	Trt	0.003785	0.116455	1			
2	1/4/2017	5/26/2018	0	Trt	0.009687	1.398635	2			
3	1/10/2017	3/15/2017	0	Cntrl	0.025288	0.20252	3			
4	1/11/2017	2/17/2017	0	Cntrl	0.029976	0.1294				=
5	1/18/2017	4/14/2017	0	Trt	0.046929	0.282283				
6	1/18/2017	7/23/2018	0	Trt	0.048199	1.557464				
7	1/20/2017	3/9/2017	0	Cntrl	0.054407	0.185398				
8	1/21/2017	6/1/2017	0	Trt	0.05543	0.416412				
9	1/21/2017	3/3/2017	0	Trt	0.057502	0.167194				
10	1/22/2017	1/26/2017	0	Cntrl	0.057688	0.069141				
11	1/22/2017	12/20/2017	0	Cntrl	0.059322	0.969148				
12	1/23/2017	2/2/2018	0	Trt	0.062278	1.088534				
13	1/24/2017	2/4/2018	0	Cntrl	0.064694	1.095082				
14	1/25/2017	5/17/2018	0	Cntrl	0.067065	1.373355				
15	1/27/2017	5/1/2018	0	Trt	0.073059	1.329166				
16	1/28/2017	2/28/2018	0	Trt	0.074532	1.159621				
17	1/28/2017	2/7/2017	0	Trt	0.075274	0.103676				
18	1/29/2017	12/23/2018	0	Trt	0.076946	1.977546				
19	2/3/2017	6/12/2017	0	Cntrl	0.091545	0.44497				
20	2/5/2017	1/12/2018	0	Trt	0.096764	1.030805				

For this data set, start dates and end dates are converted to start and end times through a transformation.

The transformation for converting to year times, is

$$Time = \frac{StartDate - Time0DateValue}{365}$$

For this dataset, the study began on January 1, 2017. The corresponding day value for January 1, 2017 is 42736. This value is found by entering the date in an empty column, changing the Data Type to Date & Time, and then changing the Data Type back to General. Thus, the formula is

$$Start Time = \frac{StartDate - 42736}{365}$$

and

$$End Time = \frac{EndDate - 42736}{365}$$

The units for the Start, End, and StageTimes columns are years. Any units may be used, but the units must be consistent throughout the use of the procedure. For example, if years are the units, then hazard rates must be yearly hazard rates. Conversions may be accomplished through the Survival Parameter Conversion Tool, which is available through the Tools menu.

If an individual has not had an event at the time of the stage analysis, but is still in the study, the end time should be left blank, as in

	StartDate	EndDate	Censor	Group	<i>≦∫<sub>x</sub></i> Start	<i>≩f<sub>x</sub></i> End	StageTimes	C8	C9	<b>C</b>
496	6/3/2019		0	Trt	2.421438					
497	6/8/2019		0	Cntrl	2.435071					
498	6/11/2019	8/3/2019	0	Cntrl	2.442833	2.588857				
499	6/12/2019	12/22/2019	0	Trt	2.444262	2.973864				
500	6/17/2019		0	Trt	2.460232					
501	6/19/2019	10/31/2019	0	Cntrl	2.463922	2.831271				
502	6/20/2019	10/17/2019	0	Trt	2.46738	2.793332				
503	6/21/2019	7/26/2019	0	Cntrl	2.470428	2.567093				
504	6/21/2019	7/29/2019	0	Cntrl	2.470458	2.573955				
505	6/30/2019		0	Cntrl	2.49507					
506	7/2/2019	11/25/2019	0	Cntrl	2.498895	2.900524				
507	7/2/2019	8/9/2019	1	Cntrl	2.498976	2.603059				
508	7/2/2019	9/23/2019	0	Trt	2.499687	2.727671				
509	7/2/2019	7/16/2019	0	Trt	2.500937	2.53834				
510	7/3/2019	9/16/2019	1	Trt	2.502371	2.709578				
511	7/4/2019	7/30/2019	0	Trt	2.506072	2.576769				
512	7/10/2019		0	Trt	2.522188					
513	7/11/2019		0	Trt	2.523956					
514	7/11/2019	12/21/2019	0	Trt	2.525425	2.972529				
515	7/12/2019		0	Trt	2.527532					
516	7/12/2019		0	Cntrl	2.527868					
517	7/15/2019		0	Trt	2.536789					
518	7/16/2019	8/18/2019	0	Cntrl	2.537412	2.629215				
519	7/24/2019	8/22/2019	0	Trt	2.561504	2.640745				
520	7/25/2019	12/13/2019	0	Cntrl	2.561729	2.948912				
521	7/25/2019	8/13/2019	0	Cntrl	2.563042	2.61626				
522	7/26/2019	8/3/2019	0	Cntrl	2.56695	2.587287				
523	8/2/2019		0	Trt	2.586078					
524	8/4/2019		0	Trt	2 590614					

# Example 1 – Group-Sequential Analysis

A colorectal cancer study is to be conducted to determine whether a new treatment following tumor excision will result in a tumor recurrence time that is lower than that of the current standard treatment by at least 0.2. Thus, the desired margin of superiority is 0.2. The response for each patient is time, in years, before recurrence. A one-sided test with alpha equal to 0.025 is used. The MLE Z-Test for comparing two hazard rates will be used.

The new treatment is assigned to Group 1, and the standard is assigned to Group 2, so that the null and alternative hypotheses are

$$H_0: h_1 - h_2 = -0.2$$
 ( $H_0: h_{New} = h_{Std} - 0.2$ )

versus

$$H_a: h_1 - h_2 < -0.2$$
  $(H_a: h_{New} < h_{Std} - 0.2)$ 

The design calls for five stages of one year each, if the final stage is reached. The current stage is the 3<sup>rd</sup> stage. In the design phase, a needed power of 0.90 called for 490 patients per group if the final stage is reached, based on assumed hazard rates of 0.85 and 1.3 for the new and standard treatments, respectively. Both efficacy and non-binding futility boundaries are implemented. The efficacy (alpha-spending) spending function used is the O'Brien-Fleming analog. The Hwang-Shih-DeCani (Gamma) beta-spending function with gamma parameter 1.5 is used for futility. Accrual is intended to be steady over the 5-year period of the study. Loss hazard rates of 0.03 for both groups are anticipated.

The boundary plot for stage 2 appeared as

![](_page_25_Figure_11.jpeg)

resulting in continuance of the study to stage 3.

# Setup

To run this example, complete the following steps:

#### 1 Open the GS Survival SM example dataset

- From the File menu of the NCSS Data window, select **Open Example Data**.
- Select **GS Survival SM** and click **OK**.
- 2 Specify the Group-Sequential Superiority by a Margin Analysis for Two Hazard Rates procedure options
  - Find and open the **Group-Sequential Superiority by a Margin Analysis for Two Hazard Rates** procedure using the menus or the Procedure Navigator.
  - The settings for this example are listed below and are stored in the Example 1 settings file. To load
    these settings to the procedure window, click Open Example Settings File in the Help Center or File
    menu.

Start Time Variable	Start
End Time Variable	End
Censor Indicator Variable	Censor
Censored	1
Not Censored	0
Group Variable	Group
Group 1 Value	Trt
Group 2 Value	Cntrl
Stage Times Variable	StageTimes
Current Stage (k)	3
Maximum Number of Stages (K)	5
Time Proportion at each Stage	Equally incremented
N1	490
N2	490
h1	0.85
h2	1.3
T0 (Accrual or Recruitment Time)	5
Accrual Parameter Entry	Enter Accrual Parameter Directly
Accrual Parameter	0
T (Total Time)	5
L1 (Loss Hazard Rate of Group 1)	0.03
L2 (Loss Hazard Rate of Group 2)	0.03
Superiority Margin (SM)	0.2
Future Stage Time Adjustment	Keep original design time proportions exactly
Boundaries Used	Efficacy with Futility
Hypothesis Direction	Ha: h1 - h2 < - SM  (Lower hazard rates are better)
Boundary Specification	Spending Function Calculation
Alpha	0.025
Alpha Spending Function	O'Brien-Fleming Analog
Skipped Efficacy Stages	<empty></empty>

#### Reports Tab

Reports for Current Stage	
All Reports	Checked
Confidence Level	95
Number of Simulations	2000
All Three Boxes under Conditional and Predictive Power and Boundary Crossing Probability Parameters	Checked
h1	1.1
h2	<b>1.1</b>
Planning Stage (Stage 0) Reports	
Summary using Z Scale	Checked
Plots	Checked

#### Plots Tab

Z-Statistic vs Information	Checked
Z-Statistic vs Time	Checked

Note: You can add the stage number to the plot by checking the **small box on the Plot Format button** (under Z-Statistic vs Information and Z-Statistic vs Time). When the procedure is run, select the **Stage Notes** tab and press the **Add Stage Number** button.

#### 3 Run the procedure

• Click the **Run** button to perform the calculations and generate the output.

### **Run Summary Report**

This report can be used to confirm that the input was processed as intended.

ltom	Valuo
	Value
Maximum Number of Stages (Design):	5
Current Stage:	3
Alternative Hypothesis:	h1 - h2 < - SM
Superiority Margin (SM):	0.2
Alpha Spending Function:	O'Brien-Fleming Analog
Beta Spending Function:	Hwang-Shih-DeCani (γ = 1.5)
Futility Boundaries:	Non-Binding
Alpha:	0.0250
Maximum Information:	163.3840
Z Statistic:	MLE

# Z-Values and Boundaries at Stage 3

This section gives the Z-test values and boundaries, numerically. These values are reflected in the groupsequential boundary plot. The Decision column indicates whether a boundary was crossed at each stage.

#### Z-Values and Boundaries at Stage 3

Maximum Information:	163.3840
Alternative Hypothesis:	h1 - h2 < - SM
Superiority Margin (SM):	0 2
Futility Boundaries:	Non-Binding
Z Statistic:	MLE

	7 7 (	Bound	daries			Decision	
Stage	Z-Test Value	Efficacy	Futility	Time	Proportion		
1	-1.3832	-6.5464	0.8096	1.0	0.1116	Continue	
2	-2.2761	-3.5232	-0.5204	2.0	0.3664	Continue	
3	-3.7574	-2.6445	-1.2038	3.0	0.6118	Crossed Efficacy	
4		-2.3780	-1.4633	4.0	0.7565	,	
5		-2.0183	-2.0183	5.0	1.0000		

### **Group-Sequential Boundary Plot at Stage 3**

This plot shows the Z-test values and boundaries. The efficacy boundary is crossed at Stage 3.

![](_page_28_Figure_10.jpeg)

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![](_page_29_Figure_2.jpeg)

# P-Values and Boundaries at Stage 3

This section reflects the conversion of the Z-test values and boundaries to the corresponding P-values and P-value boundaries.

P-Value	s and Bound	aries at Stag	e 3				
Maximu Alternati Superior Futility E Z Statist	m Information ive Hypothesis rity Margin (SM Boundaries: iic:	: 163.3840 s: h1 - h2 < M): 0.2 Non-Bind MLE	0 < - SM  ding				
Boundaries				Information			
Stage	P-Value	Efficacy	Futility	Time	Proportion	Decision	
1	0.08330	0.00000	0.79091	1.0	0.1116	Continue	
2	0.01142	0.00021	0.30139	2.0	0.3664	Continue	
3	0.00009	0.00409	0.11434	3.0	0.6118	Crossed Efficacy	
4		0.00870	0.07170	4.0	0.7565	-	
-		0.00170	0.00470	<b>F</b> 0	1 0000		

P-values and P-value boundaries are one-sided values.

### Current Stage Results Adjusted for Sequential Analysis (Stage 3)

This section gives appropriate adjustments to the raw results to reflect the group-sequential nature of the analysis. Additional explanation is given in early sections of this chapter: Adjusted Confidence Intervals, Adjusted p-values, and Adjusted Hazard Rate Difference.

		Group-Sequ	ential Adjust	ed Difference	Caref Laval
	Actual Difference	95.0% C	I. of Diff.	<u> </u>	Where Upper Boundary
Stage	h1 - h2 + SM	Lower	Upper	Midpoint	Is Zero
3	-0.37581	-0.72589	-0.21214	-0.47201	99.944%

The adjustments are based only on the efficacy boundaries. The futility boundaries are not used in these calculations.

### **Descriptive Statistics up to Stage 3**

This section gives the raw sample statistics for the cumulative data at each stage.

#### Descriptive Statistics up to Stage 3

	Samp	le Size	Ev Co	ent unt	Sar Hazaro	nple d Rates	Differe	ence
Stage	N1	N2	E1	E2	h1	h2	Difference	SE(Diff)
1	107	92	28	38	0.68176	1.20573	-0.52397	0.23422
2	209	195	96	116	0.68007	1.17425	-0.49418	0.12925
3	318	293	177	212	0.67756	1.25338	-0.57581	0.10002

### Information Report at Stage 3

This section gives the target and achieved information for each stage, as well as the sample sizes and hazard rates used to calculate those informations.

#### Information Report at Stage 3

Maximu Alterna Superio Alpha:	um Information: tive Hypothesis prity Margin (SN	163.3840 :: h1 - h2 < - 1): 0.2 0.0250	SM					
Stage	Target Information Proportion	Achieved Information Proportion	Target Information	Achieved Information	N1	N2	h1	h2
1	0.0983	0.1116	16.0624	18.2290	107.00	92.00	0.68176	1.20573
2	0.2947	0.3664	48.1453	59.8641	209.00	195.00	0.68007	1.17425
3	0.5220	0.6118	85.2925	99.9612	318.00	293.00	0.67756	1.25338
4	0.7594	*0.7565	124.0695	*123.6070	*335.35	*335.35	*0.67756	*1.25338
5	1.0000	*1.0000	163.3840	*163.3840	*419.19	*419.19	*0.67756	*1.25338

\* Projected value

# Alpha Spending at Stage 3

This section shows how alpha was spent (or is anticipated to be spent) across the stages.

#### Alpha Spending at Stage 3

Target Final Stage Alpha: Spending Function:		0.0250 O'Brien-Flemin				
Stage	Information Proportion	Alpha Spent this Stage	Cumulative Alpha Spent	Nominal (Boundary) Alpha	Percentage Alpha Spent this Stage	Cumulative Percentage Alpha Spent
1	0.1116	0.0000	0.0000	0.000000	0.0	0.0
2	0.3664	0.0002	0.0002	0.000213	0.9	0.9
3	0.6118	0.0039	0.0042	0.004091	15.8	16.7
4 *	0.7565	0.0058	0.0100	0.008703	23.2	39.9
5 *	1.0000	0.0150	0.0250	0.021780	60.1	100.0

\* Projected

# Beta Spending for Futility at Stage 3

This section shows how beta was spent (or is anticipated to be spent) across the stages.

Dota oponanig for Fatinty at otago o	Beta S	Spending	for	Futility	at	Stage 3
--------------------------------------	--------	----------	-----	----------	----	---------

Target Cumulative Beta at Final Stage:0.1000Spending Function for Futility:Hwang-Shih-DeCani (γ = 1.5)										
Stage	Information Proportion	Beta Spent this Stage	Cumulative Beta Spent	Nominal (Boundary) Beta	Percentage Beta Spent this Stage	Cumulative Percentage Beta Spent				
1	0.1116	0.0198	0.0198	0.790913	19.8	19.8				
2	0.3664	0.0346	0.0544	0.301391	34.6	54.4				
3	0.6118	0.0229	0.0773	0.114336	22.9	77.3				
4 *	0.7565	0.0100	0.0873	0.071699	10.0	87.3				
5 *	1.0000	0.0127	0.1000	0.021780	12.7	100.0				

\* Projected

### Conditional and Predictive Power Report at Stage 3

This section gives the conditional power for the various differences assumed. It also gives the predictive power.

#### **Conditional and Predictive Power Report at Stage 3**

Maximum Informatic Proportion Predictive	laximum Information: Iformation this Stage: roportion of Maximum Information: redictive Power: δ Conditiona lame Value Powe resign -0.4500 0.9998		163.3840 99.9612 0.6118 0.9998
δ	δ	Conditional	
Name	Value	Power	
Design	-0.4500	0.9998	_
Data	-0.5758	1.0000	
δ1	0.0000	0.4915	

The conditional power values are calculated based on Jennison and Turnbull (2000), pages 205 to 208. They do not account for future interim stages, nor futility boundaries, if applicable.

### Boundary Probabilities for $\delta = -0.45$

Using simulation based on the specified hazard rates, this section gives the estimated probabilities of crossing each of the future boundaries. Values given here will vary for each simulation.

#### Boundary Probabilities for $\delta = -0.45$

Number of Simulations:	2000
User-Entered Random Seed:	5758403
Futility Boundaries:	Non-Binding
After Efficacy Boundary Crossing:	Hold Out
After Non-Binding Futility Boundary Crossing:	Leave In
Alternative Hypothesis:	h1 - h2 < - SM
Superiority Margin (SM):	0.2
Z Statistic:	MLE
h1:	0.85
h2:	1.3
δ:	-0.45

			7 Teet	Effi	сасу	Futility		
Stage	N1	N2	Value	Boundary	Probability	Boundary	Probability	
1	107.00	92.00	-1.3832	-6.5464		0.8096		
2	209.00	195.00	-2.2761	-3.5232		-0.5204		
3	318.00	293.00	-3.7574	-2.6445		-1.2038		
4	*335.35	*335.35		-2.3780	0.9985	-1.4633	0.0000	
5	*419.19	*419.19		-2.0183	0.0015	-2.0183	0.0010	

\* Simulation sample size (Non-integer sample sizes were rounded to the next highest integer.)

### Event Summary for $\delta$ = -0.45

Using simulation based on the specified hazard rates, this section gives the estimated number of events at each future stage. Values given here will vary for each simulation.

#### Event Summary for $\delta = -0.45$

Number of Simulations:			2000			
User-Entered Random Seed:			5758403			
h1: h2:			0.85			
			1.3			
δ:			-0.45			
			7.74	Averaç Numb	ge Cumulative per of Events	
Stage	N1	N2	2 Val	ue E	1 E2	Time
1	107	92	-1.38	32 2	8 38	1.00
2	209	195	-2.27	'61 9	6 116	2.00
3	318	293	-3.75	574 17	7 212	3.00

\* Simulation sample size (Non-integer sample sizes were rounded to the next highest integer.)

\*335.35

\*419.19

\*\* Simulation average

\*335.35

\*419.19

4

5

\*\*293.58

\*\*361.04

\*4.00

\*5.00

\*\*269.38

\*\*336.60

# Boundary Probabilities for $\delta$ = -0.5758126

#### Boundary Probabilities for $\delta = -0.5758126$

Number of Simulations:	2000
User-Entered Random Seed:	5758403
Futility Boundaries:	Non-Binding
After Efficacy Boundary Crossing:	Hold Out
After Non-Binding Futility Boundary Crossing:	Leave In
Alternative Hypothesis:	h1 - h2 < - SM
Superiority Margin (SM):	0.2
Z Statistic:	MLE
h1:	0.6775628
h2:	1.253375
δ:	-0.5758126

			7 To at	Effi	сасу	Futility		
Stage	N1	N2	Value	Boundary	Probability	Boundary	Probability	
1	107.00	92.00	-1.3832	-6.5464		0.8096		
2	209.00	195.00	-2.2761	-3.5232		-0.5204		
3	318.00	293.00	-3.7574	-2.6445		-1.2038		
4	*335.35	*335.35		-2.3780	1.0000	-1.4633	0.0000	
5	*419.19	*419.19		-2.0183	0.0000	-2.0183	0.0000	

\* Simulation sample size (Non-integer sample sizes were rounded to the next highest integer.)

# Event Summary for $\delta$ = -0.5758126

#### Event Summary for $\delta = -0.5758126$

		Average Cumulative	
δ:	-0.5758126		
h2:	1.253375		
h1:	0.6775628		
User-Entered Random Seed:	5758403		
Number of Simulations:	2000		

			Number of Events			
Stage	N1	N2	Z-Test Value	E1	E2	Time
1	107	92	-1.3832	28	38	1.00
2	209	195	-2.2761	96	116	2.00
3	318	293	-3.7574	177	212	3.00
4	*335.35	*335.35		**256.29	**292.08	*4.00
5	*419.19	*419.19		**319.88	**359.06	*5.00

\* Simulation sample size (Non-integer sample sizes were rounded to the next highest integer.) \*\* Simulation average

# Boundary Probabilities for $\delta = 0$

#### Boundary Probabilities for $\delta = 0$

Number of Simulations:	2000
User-Entered Random Seed:	5758403
Futility Boundaries:	Non-Binding
After Efficacy Boundary Crossing:	Hold Out
After Non-Binding Futility Boundary Crossing:	Leave In
Alternative Hypothesis:	h1 - h2 < - SM
Superiority Margin (SM):	0.2
Z Statistic:	MLE
h1:	1.1
h2:	1.1
δ:	0

			7 Teet	Efficacy		Futility	
Stage	N1	N2	Z-Test Value	Boundary	Probability	Boundary	Probability
1	107.00	92.00	-1.3832	-6.5464		0.8096	
2	209.00	195.00	-2.2761	-3.5232		-0.5204	
3	318.00	293.00	-3.7574	-2.6445		-1.2038	
4	*335.35	*335.35		-2.3780	0.6210	-1.4633	0.0205
5	*419.19	*419.19		-2.0183	0.0435	-2.0183	0.5575

\* Simulation sample size (Non-integer sample sizes were rounded to the next highest integer.)

# Event Summary for $\delta = 0$

#### Event Summary for $\delta = 0$

Number of Simulations:	2000			
User-Entered Random Seed:	5758403			
h1:	1.1			
h2:	1.1			
δ:	0			
Average Cumulative				

				Number	of Events	
Stage	N1	N2	Z-Test Value	E1	E2	Time
1	107	92	-1.3832	28	38	1.00
2	209	195	-2.2761	96	116	2.00
3	318	293	-3.7574	177	212	3.00
4	*335.35	*335.35		**285.34	**286.24	*4.00
5	*419.19	*419.19		**353.73	**351.98	*5.00

\* Simulation sample size (Non-integer sample sizes were rounded to the next highest integer.) \*\* Simulation average

### Z-Values and Boundaries at Stage 0

This section gives the boundaries that were initially projected at the planning stage.

#### Z-Values and Boundaries at Stage 0

Maximum Information: Alternative Hypothesis: Superiority Margin (SM):	163.3840 h1 - h2 < - SM  0.2
Futility Boundaries:	Non-Binding
Z Statistic:	MLE

7 Tool		Boundaries				
Stage	Z-Test Value	Efficacy	Futility	Time	Proportion	Decision
1			0.9347	1.0	0.0983	
2		-3.9670	-0.1997	2.0	0.2947	
3		-2.8930	-0.9507	3.0	0.5220	
4		-2.3456	-1.5189	4.0	0.7594	
5		-2.0170	-2.0170	5.0	1.0000	

## Group-Sequential Boundary Plot at Planning Stage (Stage 0)

This plot shows the projected boundaries at the planning stage.

![](_page_36_Figure_10.jpeg)

![](_page_36_Figure_11.jpeg)

![](_page_37_Figure_2.jpeg)

### **Kaplan-Meier Survival Plot**

In order to obtain a Kaplan-Meier survival curve plot, the data must be converted into a form that can be used in the Kaplan-Meier Curves (Logrank Tests) procedure:

- 1. Rows with blank end times should be identified as censored.
- 2. Blank end times must be filled in with the current stage time.
- 3. An Elapsed Time column should be created by subtracting start times from end times.

When these steps are taken, the GS Survival SM dataset becomes the GS Survival SM B dataset.

### Setup

To run this example, complete the following steps:

#### 1 Open the GS Survival SM B example dataset

- From the File menu of the NCSS Data window, select **Open Example Data**.
- Select GS Survival SM B and click OK.

#### 2 Specify the Kaplan-Meier Curves (Logrank Tests) procedure options

• Find and open the **Kaplan-Meier Curves (Logrank Tests)** procedure using the menus or the Procedure Navigator.

• The settings for this example are listed below and are stored in the **Example GS** settings file. To load these settings to the procedure window, click **Open Example Settings File** in the Help Center or File menu.

Variables Tab
(Elapsed) Time VariableElapsed
Censor Variable
Failed0
Censored1
Group VariableGroup
Reports Tab
Data Summary
All Other Reports Unchecked
Plots Tab
Individual-Group PlotsUnchecked
Combined Plot(s)Checked
Kaplan-Meier Survival/Reliability Plot Format (Click the Button)
Kaplan-Meier Survival LineChecked
Confidence Limits
Confidence Limits - Fill Checked

#### 3 Run the procedure

• Click the **Run** button to perform the calculations and generate the output.

### Kaplan-Meier Survival Curve(s)

This plot shows the survival curves for the two groups.

![](_page_38_Figure_9.jpeg)

![](_page_38_Figure_10.jpeg)

# Example 2 – Skipping Stage Boundaries

Suppose that the setup is the same as in Example 1, except that the first two futility boundaries are skipped.

### Setup

To run this example, complete the following steps:

#### 1 Open the GS Survival SM example dataset

- From the File menu of the NCSS Data window, select Open Example Data.
- Select **GS Survival SM** and click **OK**.
- 2 Specify the Group-Sequential Superiority by a Margin Analysis for Two Hazard Rates procedure options
  - Find and open the **Group-Sequential Superiority by a Margin Analysis for Two Hazard Rates** procedure using the menus or the Procedure Navigator.
  - The settings for this example are listed below and are stored in the **Example 2** settings file. To load these settings to the procedure window, click **Open Example Settings File** in the Help Center or File menu.

Variables Tab	
Skipped Futility Stages	

#### 3 Run the procedure

• Click the **Run** button to perform the calculations and generate the output.

### **Run Summary Report**

The skipped futility stages are now reported.

#### Run Summary Report

Item	Value
Maximum Number of Stages (Design):	5
Skipped Futility Stage(s):	12
Current Stage:	3
Alternative Hypothesis:	h1 - h2 < - SM
Superiority Margin (SM):	0.2
Alpha Spending Function:	O'Brien-Fleming Analog
Beta Spending Function:	Hwang-Shih-DeCani (γ = 1.5)
Futility Boundaries:	Non-Binding
Alpha:	0.0250
Maximum Information:	163.3840
Z Statistic:	MLE

### Z-Values and Boundaries at Stage 3

The futility boundaries change slightly from those where no boundaries are skipped.

#### Z-Values and Boundaries at Stage 3

Maximum Information:	163.3840
Alternative Hypothesis:	h1 - h2 < - SM
Superiority Margin (SM):	0.2
Futility Boundaries:	Non-Binding
Z Statistic:	MI F
Z Statistic:	MLE
Futility Boundaries:	Non-Binding
Z Statistic:	MLE

	7 7(	Bound	daries			
Stage	Z-Test Value	Efficacy	Futility	Time	Proportion	Decision
1	-1.3832	-6.5464		1.0	0.1116	Continue
2	-2.2761	-3.5232		2.0	0.3664	Continue
3	-3.7574	-2.6445	-1.4412	3.0	0.6118	Crossed Efficacy
4		-2.3780	-1.5233	4.0	0.7565	
5		-2.0183	-2.0183	5.0	1.0000	

# **Group-Sequential Boundary Plot at Stage 3**

The plot now has two futility boundaries skipped.

![](_page_40_Figure_10.jpeg)

#### Group-Sequential Boundary Plot at Stage 3

![](_page_41_Figure_2.jpeg)

# Example 3 – Sample Size Re-estimation and Boundary Probabilities

Suppose that the setup is the same as in Example 1, except that only the data up through Stage 2 has been collected.

### Setup

To run this example, complete the following steps:

#### 1 Open the GS Survival SM 2 Stages example dataset

- From the File menu of the NCSS Data window, select **Open Example Data**.
- Select GS Survival SM 2 Stages and click OK.
- 2 Specify the Group-Sequential Superiority by a Margin Analysis for Two Hazard Rates procedure options
  - Find and open the **Group-Sequential Superiority by a Margin Analysis for Two Hazard Rates** procedure using the menus or the Procedure Navigator.
  - The settings for this example are listed below and are stored in the **Example 3** settings file. To load these settings to the procedure window, click **Open Example Settings File** in the Help Center or File menu.

Variables Tab	
Current Stage (k)	.2

#### 3 Run the procedure

• Click the **Run** button to perform the calculations and generate the output.

### **Run Summary Report**

This report can be used to confirm that the input was processed as intended.

#### Run Summary Report

ltem	Value
Maximum Number of Stages (Design):	5
Current Stage:	2
Alternative Hypothesis:	h1 - h2 < - SM
Superiority Margin (SM):	0.2
Alpha Spending Function:	O'Brien-Fleming Analog
Beta Spending Function:	Hwang-Shih-DeCani ( $\gamma = 1.5$ )
Futility Boundaries:	Non-Binding
Alpha:	0.0250
Maximum Information:	163.3840
Z Statistic:	MLE

## Z-Values and Boundaries at Stage 2

Examining the Z-Test values, the boundaries, and the decisions, no boundary has been crossed at this stage.

#### Z-Values and Boundaries at Stage 2

163.3840 h1 - h2 < - SM  0.2 Non-Binding MI F
MLE

		Bound	daries			
Stage	Z-Test Value	Efficacy	Futility	Time	Information Proportion	Decision
1	-1.3832	-6.5464	0.8114	1.0	0.1116	Continue
2	-2.2761	-3.5232	-0.5170	2.0	0.3664	Continue
3		-2.9318	-0.8714	3.0	0.5132	
4		-2.3543	-1.5050	4.0	0.7541	
5		-2.0155	-2.0155	5.0	1.0000	

## Group-Sequential Boundary Plot at Stage 2

This plot shows the Z-test values in the vicinity of the efficacy boundary side.

![](_page_43_Figure_10.jpeg)

#### Group-Sequential Boundary Plot at Stage 2

![](_page_44_Figure_2.jpeg)

# Descriptive Statistics up to Stage 2

#### Descriptive Statistics up to Stage 2

	Samp	le Size	E\ Co	vent ount	San Hazaro	nple I Rates	Differe	ence
Stage	N1	N2	E1	E2	h1	h2	Difference	SE(Diff)
1 2	107 209	92 195	28 96	38 116	0.68176 0.68007	1.20573 1.17425	-0.52397 -0.49418	0.23422 0.12925

### Information Report at Stage 2 (Gives Sample Size Re-estimation)

This section shows that the target sample size for the next stage should be 232 per group.

#### Information Report at Stage 2

Maximum Information:	163.3840
Alternative Hypothesis:	h1 - h2 < - SM
Superiority Margin (SM):	0.2
Alpha:	0.0250
Target	Achieved

Stage	Information Proportion	Information Proportion	Target Information	Achieved Information	N1	N2	h1	h2
1	0.0983	0.1116	16.0624	18.2290	107.00	92.00	0.68176	1.20573
2	0.2947	0.3664	48.1453	59.8641	209.00	195.00	0.68007	1.17425
3	0.5220	*0.5132	85.2925	*83.8437	*231.32	*231.32	*0.68007	*1.17425
4	0.7594	*0.7541	124.0695	*123.2136	*308.42	*308.42	*0.68007	*1.17425
5	1.0000	*1.0000	163.3840	*163.3840	*385.53	*385.53	*0.68007	*1.17425

\* Projected value

# Alpha Spending at Stage 2

This section shows how alpha was spent (or is anticipated to be spent) across the stages.

#### Alpha Spending at Stage 2

Target F Spendin	inal Stage Alpha: g Function:	0.0250 O'Brien-Flemin	g Analog			
Stage	Information Proportion	Alpha Spent this Stage	Cumulative Alpha Spent	Nominal (Boundary) Alpha	Percentage Alpha Spent this Stage	Cumulative Percentage Alpha Spent
1	0.1116	0.0000	0.0000	0.000000	0.0	0.0
2	0.3664	0.0002	0.0002	0.000213	0.9	0.9
3 *	0.5132	0.0015	0.0018	0.001685	6.2	7.0
4 *	0.7541	0.0081	0.0099	0.009280	32.4	39.4
5 *	1.0000	0.0151	0.0250	0.021924	60.6	100.0

\* Projected

## Beta Spending for Futility at Stage 2

This section shows how beta was spent (or is anticipated to be spent) across the stages.

beta openang for rating at otage z	Beta	Spending	for	Futility	at	Stage 2
------------------------------------	------	----------	-----	----------	----	---------

Target C Spendin	Cumulative Beta at g Function for Fut	t Final Stage: 0 ility: H	.1000 wang-Shih-DeCa	ni (γ = 1.5)		
Stage	Information Proportion	Beta Spent this Stage	Cumulative Beta Spent	Nominal (Boundary) Beta	Percentage Beta Spent this Stage	Cumulative Percentage Beta Spent
1	0.1116	0.0198	0.0198	0.791446	19.8	19.8
2	0.3664	0.0346	0.0544	0.302563	34.6	54.4
3 *	0.5132	0.0147	0.0691	0.191765	14.7	69.1
4 *	0.7541	0.0181	0.0872	0.066163	18.1	87.2
5 *	1.0000	0.0128	0.1000	0.021924	12.8	100.0

\* Projected

### Conditional and Predictive Power Report at Stage 2

Conditional power and predictive power are described earlier in the Technical details section. The predictive power does not depend on an assumed difference, as does the conditional power.

#### **Conditional and Predictive Power Report at Stage 2**

Maximum Information Proportion Predictive	n Information on this Stage n of Maximur e Power:	: : m Information:	163.3840 59.8641 0.3664 0.9145
δ	δ	Conditional	
Name	Value	Power	
Design	-0.4500	0.9650	_
Data	-0.4942	0.9881	
δ1	0.0000	0.0028	

The conditional power values are calculated based on Jennison and Turnbull (2000), pages 205 to 208. They do not account for future interim stages, nor futility boundaries, if applicable.

### Boundary Probabilities for $\delta = -0.45$

Given the data that have already accumulated in the first two stages, and assuming hazard rates of 0.85 and 1.3 going forward, this report gives the simulation probabilities of crossing each of the future boundaries. The sum of the efficacy boundary probabilities is another estimate of the conditional power.

#### Boundary Probabilities for $\delta = -0.45$

Number of Simulations: User-Entered Random Seed: Futility Boundaries: After Efficacy Boundary Crossing: After Non-Binding Futility Boundary Crossing: Alternative Hypothesis: Superiority Margin (SM):	2000 6307504 Non-Binding Hold Out Leave In h1 - h2 < - SM  0.2
Superiority Margin (SM): Z Statistic:	0.2 MLE
h1:	0.85
h2:	1.3
δ:	-0.45

			7 Teet	Effi	icacy	Futility		
Stage	N1	N2	Value	Boundary	Probability	Boundary	Probability	
1	107.00	92.00	-1.3832	-6.5464		0.8114		
2	209.00	195.00	-2.2761	-3.5232		-0.5170		
3	*231.32	*231.32		-2.9318	0.3250	-0.8714	0.0030	
4	*308.42	*308.42		-2.3543	0.4780	-1.5050	0.0195	
5	*385.53	*385.53		-2.0155	0.1360	-2.0155	0.0670	

\* Simulation sample size (Non-integer sample sizes were rounded to the next highest integer.)

# Event Summary for $\delta$ = -0.45

Using simulation based on the specified hazard rates, this section gives the estimated number of events at each future stage. Values given here will vary for each simulation.

#### Event Summary for $\delta = -0.45$

Number of Simulations:	2000
User-Entered Random Seed:	6307504
h1:	0.85
h2:	1.3
δ:	-0.45

			7-Tost	Average C Number	Cumulative of Events	
Stage	N1	N2	Value	E1	E2	Time
1	107	92	-1.3832	28	38	1.00
2	209	195	-2.2761	96	116	2.00
3	*231.32	*231.32		**177.78	**197.84	*3.00
4	*308.42	*308.42		**245.29	**265.12	*4.00
5	*385.53	*385.53		**306.52	**328.60	*5.00

\* Simulation sample size (Non-integer sample sizes were rounded to the next highest integer.)

\*\* Simulation average

### Boundary Probabilities for $\delta$ = -0.4941798

Given the data that have already accumulated in the first two stages, and assuming hazard rates of 0.6800724 and 1.174252 going forward, this report gives the simulation probabilities of crossing each of the future boundaries.

#### Boundary Probabilities for $\delta$ = -0.4941798

Number of Simulations:	2000
User-Entered Random Seed:	6307504
Futility Boundaries:	Non-Binding
After Efficacy Boundary Crossing:	Hold Out
After Non-Binding Futility Boundary Crossing:	Leave In
Alternative Hypothesis:	h1 - h2 < - SM
Superiority Margin (SM):	0.2
Z Statistic:	MLE
h1:	0.6800724
h2:	1.174252
δ:	-0.4941798

<b>Stage</b> 1 2 3				Effi	icacy	Futility		
	N1	N2	Value	Boundary	Probability	Boundary	Probability	
1	107.00	92.00	-1.3832	-6.5464		0.8114		
2	209.00	195.00	-2.2761	-3.5232		-0.5170		
3	*231.32	*231.32		-2.9318	0.5075	-0.8714	0.0005	
4	*308.42	*308.42		-2.3543	0.4295	-1.5050	0.0025	
5	*385.53	*385.53		-2.0155	0.0530	-2.0155	0.0105	

\* Simulation sample size (Non-integer sample sizes were rounded to the next highest integer.)

# Event Summary for $\delta$ = -0.4941798

#### Event Summary for $\delta = -0.4941798$

Number of Simulations:	2000
User-Entered Random Seed:	6307504
h1:	0.6800724
h2:	1.174252
δ:	-0.4941798

				Average Cumulative Number of Events			
Stage	N1	N2	Z-Test Value	E1	E2	Time	
1	107	92	-1.3832	28	38	1.00	
2	209	195	-2.2761	96	116	2.00	
3	*231.32	*231.32		**166.04	**193.24	*3.00	
4	*308.42	*308.42		**229.70	**259.96	*4.00	
5	*385.53	*385.53		**289.45	**323.20	*5.00	

\* Simulation sample size (Non-integer sample sizes were rounded to the next highest integer.)

\*\* Simulation average

### Boundary Probabilities for $\delta = 0$

Given the data that have already accumulated in the first two stages, and assuming hazard rates of 1.1 and 1.1 going forward, this report gives the simulation probabilities of crossing each of the future boundaries.

#### Boundary Probabilities for $\delta = 0$

Number of Simulations:	2000
User-Entered Random Seed:	6307504
Futility Boundaries:	Non-Binding
After Efficacy Boundary Crossing:	Hold Out
After Non-Binding Futility Boundary Crossing:	Leave In
Alternative Hypothesis:	h1 - h2 < - SM
Superiority Margin (SM):	0.2
Z Statistic:	MLE
h1:	1.1
h2:	1.1
δ:	0

Stage		7 Test		Effi	сасу	Futility	
	N1	N2	Value	Boundary	Probability	Boundary	Probability
1	107.00	92.00	-1.3832	-6.5464		0.8114	
2	209.00	195.00	-2.2761	-3.5232		-0.5170	
3	*231.32	*231.32		-2.9318	0.0005	-0.8714	0.4140
4	*308.42	*308.42		-2.3543	0.0020	-1.5050	0.9400
5	*385.53	*385.53		-2.0155	0.0015	-2.0155	0.9975

\* Simulation sample size (Non-integer sample sizes were rounded to the next highest integer.)

### Event Summary for $\delta = 0$

#### Event Summary for $\delta = 0$

Number of Simulations: User-Entered Random Seed:	2000 6307504
h1:	1.1
h2:	1.1
δ:	0

			7 Teet	Average C Number	Average Cumulative Number of Events		
Stage	N1	N2	Value	E1	E2	Time	
1	107	92	-1.3832	28	38	1.00	
2	209	195	-2.2761	96	116	2.00	
3	*231.32	*231.32		**192.10	**190.25	*3.00	
4 5	*308.42 *385.53	*308.42 *385.53		**261.95 **323.56	**256.53 **319.58	*4.00 *5.00	

\* Simulation sample size (Non-integer sample sizes were rounded to the next highest integer.)

\*\* Simulation average