SURVIVAL ANALYSIS

Chapter 4. Evaluating the Proportional Hazards Assumption

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Background

Cox PH model:

$$h(t,\mathbf{X}) = h_0(t)e^{\sum_{i=1}^p \beta_i X_i}$$

$$\mathbf{X} = (X_1, X_2, \dots, X_p)$$
 explanatory/
predictor variables

$$h_0(t)$$
 \times $e^{\sum_{i=1}^{p} \beta_i X_i}$

Baseline hazard | Exponential |
Involves t but | Involves X 's but |
not X 's | not t (X 's are time-independent)

X's involving t: time-dependent

Requires extended Cox model

(no PH)

Chapter 6

- Recall from the previous chapter that the general form of the Cox PH model gives an expression for the hazard at time t for an individual with a given specification of a set of explanatory variables denoted by the bold X.
- The Cox model formula says that the hazard at time t is the product of two quantities. The first of these, $h_0(t)$, is called the **baseline** hazard function. The second quantity is the exponential expression e to the linear sum of $\beta_i X_i$, where the sum is over the p explanatory X variables.
- An important feature of this formula, which concerns the proportional hazards (PH) assumption, is that the baseline hazard is a function of t, but does not involve the X's, whereas the exponential expression involves the X's, but does not involve t. The X's here are called time-independent X's.
- It is possible, nevertheless, to consider X's that do involve t. Such X's are called **time-dependent** variables. If time-dependent variables are considered, the Cox model form may still be used, but such a model no longer satisfies the PH assumption, and is called the **extended Cox model**. We will discuss this extended Cox model in Chapter 6 of this series.

Background

Hazard ratio formula:

$$\widehat{HR} = \exp\left[\sum_{i=1}^{p} \hat{\beta}_i \left(X_i^* - X_i\right)\right]$$

where $\mathbf{X}^* = (X_1^*, X_2^*, \dots, X_p^*)$ and $\mathbf{X} = (X_1, X_2, \dots, X_p)$ denote the two sets of X's.

Adjusted survival curves: 0 or 1

Comparing
$$E$$
 groups:
$$\hat{S}(t, \mathbf{X}) = [\hat{S}_0(t)]^{\exp[\beta_1 E + \sum_{i \neq 1} \hat{\beta}_i \overline{X}_i]}$$

Single curve:

$$\hat{S}(t, \overline{\mathbf{X}})[\hat{S}_0(t)]^{\exp[\hat{\beta}_i X_i]}$$

PH assumption:

$$\frac{\hat{h}(t, \mathbf{X}^*)}{\hat{h}(t, \mathbf{X})} = \hat{\theta}$$
, constant over t

i.e.,
$$\hat{h}(t, \mathbf{X}^*) = \hat{\theta}\hat{h}(t, \mathbf{X})$$

Hazards cross \Rightarrow PH not met Hazards don't cross \Rightarrow PH met

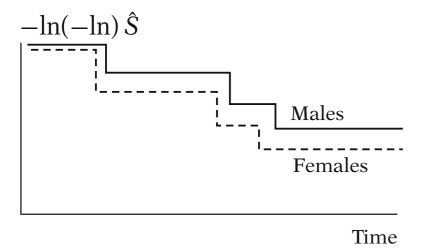
- From the Cox PH model, we can obtain a general formula, shown here, for estimating a hazard ratio that compares two specifications of the X's, defined as X^* and X.
- We can also obtain from the Cox model an expression for an adjusted survival curve. Here we show a general formula for obtaining adjusted survival curves comparing two groups adjusted for other variables in the model. Below this, we give a formula for a single adjusted survival curve that adjusts for all X's in the model. Computer packages for these formulae use the mean value of each X being adjusted in the computation of the adjusted curve.
- The Cox PH model assumes that the hazard ratio comparing any two specifications of predictors is constant over time. Equivalently, this means that the hazard for one individual is proportional to the hazard for any other individual, where the proportionality constant is independent of time.
- The PH assumption is not met if the graph of the hazards cross for two or more categories of a predictor of interest. However, even if the hazard functions do not cross, it is possible that the PH assumption is not met. Thus, rather than checking for crossing hazards, we must use other approaches to evaluate the reasonableness of the PH assumption.

Checking the Proportional Hazards Assumption: Overview

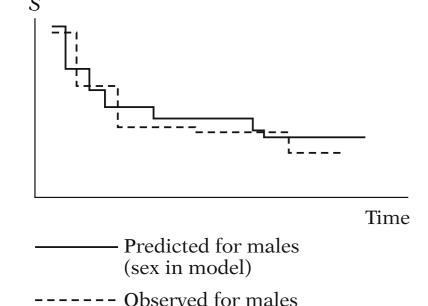
Three approaches:

- graphical
- goodness-of-fit test
- time-dependent variables

Graphical techniques: -ln(-ln) *S* curves parallel?



Observed vs. predicted: Close?



- There are three general approaches for assessing the PH assumption, again listed here. We now briefly overview each approach, starting with graphical techniques.
- There are two types of graphical techniques available. The most popular of these involves comparing estimated –In(–In) survivor curves over different (combinations of) categories of variables being investigated. We will describe such curves in detail in the next section. Parallel curves, say comparing males with females, indicate that the PH assumption is satisfied, as shown in this illustration for the variable Sex.
- An alternative graphical approach is to compare observed with predicted survivor curves. The observed curves are derived for categories of the variable being assessed, say, Sex, without putting this variable in a PH model. The predicted curves are derived with this variable included in a PH model. If observed and predicted curves are close, then the PH assumption is reasonable.

Checking the Proportional Hazards Assumption: Overview

Goodness-of-fit (GOF) tests:

- Large sample *Z* or chi-square statistics
- Gives p-value for evaluating PH assumption for each variable in the model.

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p-value large \Rightarrow PH satisfied (e.g. P > 0.10)
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p-value small \Rightarrow PH not satisfied (e.g. P < 0.05)

Time-dependent covariates:

Extended Cox model: Add product term involving some function of time. • A second approach for assessing the PH assumption involves goodness-of-fit (GOF) tests. This approach provides large sample Z or chi-square statistics which can be computed for each variable in the model, adjusted for the other variables in the model. A p-value derived from a standard normal statistic is also given for each variable. This p-value is used for evaluating the PH assumption for that variable. A nonsignificant (i.e., large) p-value, say greater than 0.10, suggest that the PH assumption is reasonable, whereas a small p-value, say less than 0.05, suggests that the variable being tested does not satisfy this assumption.

 When time-dependent variables are used to assess the PH assumption for a time-independent variable, the Cox model is extended to contain **product** (i.e., interaction) **terms** involving the time-independent variable being assessed and some function of time.

Checking the Proportional Hazards Assumption: Overview

EXAMPLE

 $h(t, X) = h_0(t) \exp[\beta \operatorname{Sex} + \delta(\operatorname{Sex} \times t)]$ $\delta \neq 0 \Rightarrow \operatorname{PH} \text{ assumption violated}$

GOF provides test statistic
Graphical: subjective
Time-dependent: computationally
cumbersome

GOF: global, may not detect specific departures from PH

- For example, if the PH assumption is being assessed for Sex, a Cox model might be extended to include the variable "Sex × t" in addition to Sex. If the coefficient of the product term turns out to be significant, we can conclude that the PH assumption is violated for Sex.
- The GOF approach provides a single test statistic for each variable being assessed. This approach is not as subjective as the graphical approach nor as cumbersome computationally as the timedependent variable approach. Nevertheless, a GOF test may be too "global" in that it may not detect specific departures from the PH assumption that may be observed from the other two approaches.

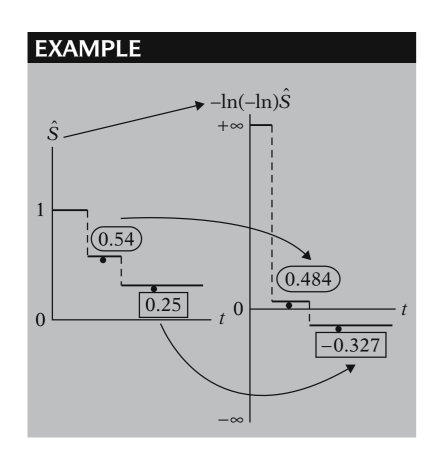
- log-log survival curves
- observed versus expected survival curves

$$\log - \log \hat{S} = \text{transformation of } \hat{S}$$

= $-\ln(-\ln \hat{S})$

- $\ln \hat{S}$ is negative $\Rightarrow -(\ln \hat{S})$ is positive.
- can't take $\log of \ln \hat{S}$, but can take $\log of (-\ln \hat{S})$.
- $-\ln(-\ln \hat{S})$ may be positive or negative.

- The two graphical approaches for checking the PH assumption are comparing log-log survival curves and comparing observed versus expected survival curves. We first explain what a -ln -ln survival curve is and how it is used.
- A log-log survival curve is simply a transformation of an estimated survival curve that results from taking the natural log of an estimated survival probability twice. Mathematically, we write a log-log curve as $-\ln(-\ln S^{\circ})$. Note that the log of a probability such as S° is always a negative number. Because we can only take logs of positive numbers, we need to negate the first log before taking the second log. The value for $-\ln(-\ln S^{\circ})$ may be positive or negative, either of which is acceptable, because we are not taking a third log.



- As an example, in the graph at left, the estimated survival probability of 0.54 is transformed to a -ln-ln value of 0.484. Similarly, the point 0.25 on the survival curve is transformed to a -ln-ln value of -0.327.
- Note that because the survival curve is usually plotted as a step function, so will the log-log curve be plotted as a step function.

EXAMPLE $\hat{S} = 0.54: \text{ want } -\ln(-\ln 0.54)$ $-\ln(-\ln 0.54) = -\ln(0.616)$ $\text{since } \ln(0.54) = -0.616$ $-\ln(0.616) = 0.484$ $\text{since } \ln(0.616) = -0.484$ Thus, $(-\ln(-\ln 0.54) = 0.484)$

• To illustrate the computation of a log-log value, suppose we start with an estimated survival probability of 0.54. Then the log-log transformation of this value is -ln(-ln 0.54), which is -ln(0.616), because ln(0.54) equals -0.616. Now, continuing further, -ln(0.616) equals 0.484, because ln(0.616) equals -0.484. Thus, the transformation -ln(-ln 0.54) equals 0.484.

ANOTHER EXAMPLE

$$\hat{S} = 0.25$$
: want $-\ln(-\ln 0.25)$
 $-\ln(-\ln 0.25) = -\ln(1.386) = -0.327$
Thus, $-\ln(-\ln 0.25) = -0.327$

• As another example, if the estimated survival probability is 0.25, then -ln(-ln 0.25) equals -ln(1.386), which equals -0.327.

y-axis scale:

$$\begin{vmatrix} 1 \\ 0 \end{vmatrix} \hat{S} \qquad \begin{vmatrix} +\infty \\ -\ln(-\ln)\hat{S} \end{vmatrix}$$

 $\log - \log \hat{S}$ for the Cox PH model:

 Note that the scale of the y-axis of an estimated survival curve ranges between 0 and 1, whereas the corresponding scale for a -ln(-ln) curve ranges between -∞ and +∞.

 We now show why the PH assumption can be assessed by evaluating whether or not log-log curves are parallel. To do this, we must first describe the log-log formula for the Cox PH model.

Cox PH hazard function:

$$h(t, \mathbf{X}) = h_0(t)e^{\sum_{j=1}^{p} \beta_j X_j}$$
From math

Cox PH survival function:

$$S(t,\mathbf{X}) = [S_0(t)]^{e^{\sum_{j=1}^{p} \beta_j X_j}}$$

 $log-log \Rightarrow takes logs twice$

log #1:

$$\ln S(t, \mathbf{X}) = e^{\sum_{i=1}^{p} \beta_i X_i} \times \ln S_0(t)$$

$$0 < S(t, \mathbf{X}) < 1$$

 $\ln(\text{probability}) = \text{negative value},$ so $\ln S(t, \mathbf{X})$ and $\ln S_0(t)$ are negative.

But $-\ln S(t, \mathbf{X})$ is positive, which allows us to take logs again.

- We start with the formula for the survival curve that corresponds to the hazard function for the Cox PH model. Recall that there is a mathematical relationship between any hazard function and its corresponding survival function. We therefore can obtain the formula shown here for the survival curve for the Cox PH model. In this formula, the expression $S_0(t)$ denotes the baseline survival function that corresponds to the baseline hazard function $h_0(t)$.
- The log-log formula requires us to take logs of this survival function twice. The first time we take logs we get the expression shown here.
- Now since $S(t, \mathbf{X})$ denotes a survival probability, its value for any t and any specification of the vector \mathbf{X} will be some number between 0 and 1. It follows that the natural log of any number between 0 and 1 is a negative number, so that the log of $S(t, \mathbf{X})$ as well as the log of $S(t, \mathbf{X})$ are both negative numbers. This is why we have to put a minus sign in front of this expression before we can take logs a second time, because there is no such thing as the log of a negative number.

log #2:

or

 $\ln[-\ln S(t, \mathbf{X})]$

$$\ln[-\ln S(t, \mathbf{X})]$$

$$= \ln \left[-e^{\sum_{i=1}^{p} \beta_{i} X_{i}} \times \ln S_{0}(t) \right]$$

$$= \ln \left[e^{\sum_{i=1}^{p} \beta_{i} X_{i}} \right] + \ln[-\ln S_{0}(t)]$$

$$= \sum_{i=1}^{p} \beta_{i} X_{i} + \ln[-\ln S_{0}(t)]$$

$$-\ln[-\ln S(t, \mathbf{X})]$$

$$= -\sum_{i=1}^{p} \beta_{i} X_{i} - \ln[-\ln S_{0}(t)]$$

 $= + \sum_{i=1}^{p} \beta_i X_i + \ln[-\ln S(t)]$

- Thus, when taking the second log, we must obtain the log of –ln $S(t, \mathbf{X})$, as shown here. After using some algebra, this expression can be rewritten as the sum of two terms, one of which is the linear sum of the $\beta_i X_i$ and the other is the log of the negative log of the baseline survival function.
- This second log may be either positive or negative, and we aren't taking any more logs, so we actually don't have to take a second negative. However, for consistency's sake, a common practice is to put a minus sign in front of the second log to obtain the -ln -ln expression shown here. Nevertheless, some software packages do not use a second minus sign.

Two individuals:

$$\mathbf{X_1} = (X_{11}, X_{12}, \dots, X_{1p})$$

 $\mathbf{X_2} = (X_{21}, X_{22}, \dots, X_{2p})$

$$\begin{cases}
-\ln[-\ln S(t, \mathbf{X}_{1})] \\
= -\sum_{i=1}^{p} \beta_{i} X_{1i} - \ln[-\ln S_{0}(t)] \\
-\ln[-\ln S(t, \mathbf{X}_{2})] \\
= -\sum_{i=1}^{p} \beta_{i} X_{2i} - \ln[-\ln S_{0}(t)]
\end{cases}$$

- Now suppose we consider two different specifications of the X vector, corresponding to two different individuals, X_1 and X_2 .
- Then the corresponding log-log curves for these individuals are given as shown here, where we have simply substituted X₁ and X₂ for X in the previous expression for the log-log curve for any individual X.

$$-\ln[-\ln S(t, \mathbf{X}_{1})]$$

$$-(-\ln[-\ln S(t, \mathbf{X}_{2})])$$

$$= \sum_{i=1}^{p} \beta_{i}(X_{2i} - X_{1i})$$

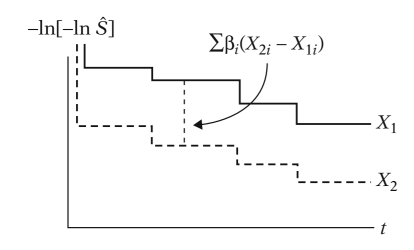
does not involve t

$$-\ln[-\ln S(t, \mathbf{X}_{1})]$$

$$= -\ln[-\ln S(t, \mathbf{X}_{2})]$$

$$+ \sum_{i=1}^{p} \beta_{i}(X_{2i} - X_{1i})$$

 Alternatively, using algebra, we can write the above equation by expressing the log-log survival curve for individual X₁ as the log-log curve for individual X₂ plus a linear sum term that is independent of t.



• The above formula says that if we use a Cox PH model and we plot the estimated log-log survival curves for individuals on the same graph, the two plots would be approximately parallel. The distance between the two curves is the linear expression involving the differences in predictor values, which does not involve time. Note, in general, if the vertical distance between two curves is constant, then the curves are parallel.

Graphical approach using log-log plots: PH model is appropriate if "empirical" plots of log-log survival curves are parallel.

Empirical plots: use $-\ln[-\ln \hat{S}]$ where

- 1. Ŝ is a KM curve
- 2. Ŝ is an adjusted survival curve for predictors satisfying the PH assumption; predictor being assessed not included in model

- The parallelism of log-log survival plots for the Cox PH model provides us with a graphical approach for assessing the PH assumption. That is, if a PH model is appropriate for a given set of predictors, one should expect that empirical plots of log-log survival curves for different individuals will be approximately parallel.
- By empirical plots, we mean plotting log-log survival curves based on Kaplan-Meier (KM) estimates that do not assume an underlying Cox model. Alternatively, one could plot log-log survival curves which have been adjusted for predictors already assumed to satisfy the PH assumption but have not included the predictor being assessed in a PH model.

EXAMPLE

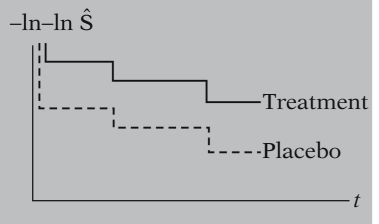
Clinical trial of leukemia patients: *T* = weeks until patient goes out of remission

Predictors (*X*'s): Rx (= 1 if placebo, 0 if treatment) log WBC

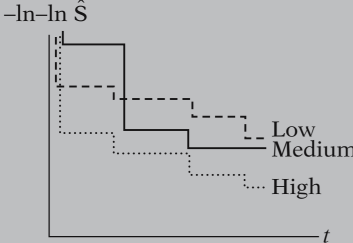
Cox PH model:

 $h(t, \mathbf{X}) = h_0(t) \exp[\beta_1 Rx + \beta_2 \log \text{WBC}]$ Assessing PH assumption: compare log-log survival curves for categories of Rx and log WBC

One-at-a-time strategy: *Rx* variable



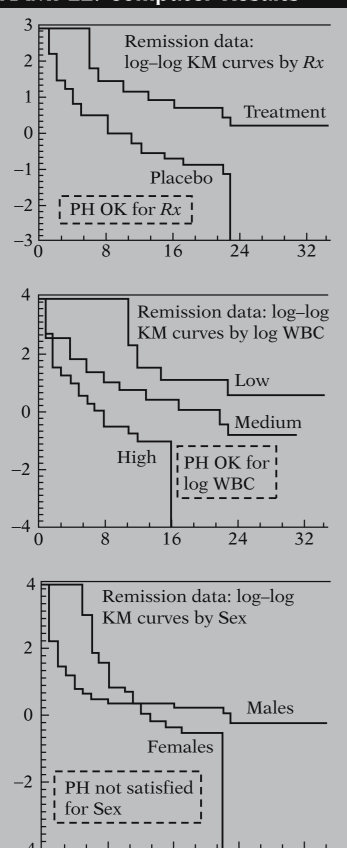
One-at-a-time strategy: log WBC



roach 1: Log-Log Plots

- As an example, suppose we consider the comparison of treatment and placebo groups in a clinical trial of leukemia patients, where survival time is time, in weeks, until a patient goes out of remission. Two predictors of interest in this study are treatment group status (1 = placebo, 0 = treatment), denoted as Rx, and log white blood cell count (log WBC), where the latter variable is being considered as a confounder.
- A Cox PH model involving both these predictors would have the form shown at the left. To assess whether the PH assumption is satisfied for either or both of these variables, we would need to compare log-log survival curves involving categories of these variables.
- One strategy to take here is to consider the variables one at a time. For the Rx variable, this amounts to plotting log-log KM curves for treatment and placebo groups and assessing parallelism. If the two curves are approximately parallel, as shown here, we would conclude that the PH assumption is satisfied for the variable Rx. If the two curves intersect or are not parallel in some other way, we would conclude that the PH assumption is not satisfied for this variable.
- For the log WBC variable, we need to categorize this variable into categories—say, low, medium, and high—and then compare plots of log-log KM curves for each of the three categories. In this illustration, the three log-log Kaplan–Meier curves are clearly nonparallel, indicating the PH assumption is not met for log WBC.

EXAMPLE: Computer Results



16

24

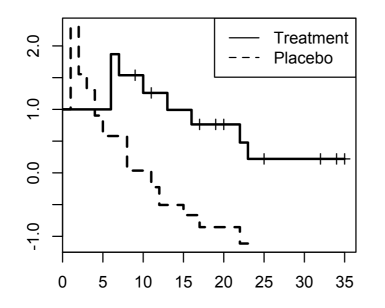
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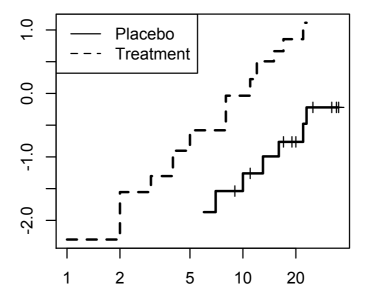
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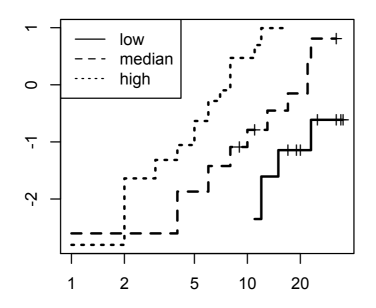
roach 1: Log-Log Plots

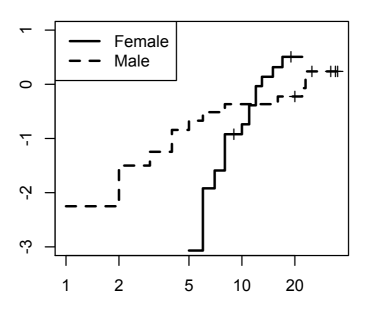
- The above examples are sketches of some of the possibilities that could occur from comparisons of log-log curves. For the actual data set containing 42 leukemia patients, computer results are shown here for each variable separately.
- We first show the log-log KM curves by treatment, Rx. Notice that the two log-log curves are roughly parallel, indicating that the Rx variable satisfies the PH assumption when being considered by itself.
- Here we show the log-log KM curves by log WBC, where we have divided this variable into low (be- low 2.3), medium (between 2.3 and 3), and high (above 3) values. Notice that there is some indication of nonparallelism below 8 days, but that overall the three curves are roughly parallel. Thus, these plots suggest that the PH assumption is more or less satisfied for the variable log WBC, when considered alone.
- As a third example, we consider the log-log KM plots categorized by Sex from the remission data. Notice that the two curves clearly intersect, and are therefore noticeably nonparallel. Thus, the variable, Sex, when considered by itself, does not appear to satisfy the PH assumption and therefore should not be incorporated directly into a Cox PH model containing the other two variables, Rx and log WBC.

```
library(survival) # loading survival package
leukemia<-read.csv('leukemia.csv',header=T)</pre>
par(mfrow=c(2,2))
res1<-survfit(Surv(Survt,Relapse)~Rx,data=leukemia)</pre>
res1.1<-res1
res1.1\surv<- -log(-log(res1\surv))
plot(res1.1, lty=1:2, lwd=2)
legend('topright',c('Treatment','Placebo'),lty=1:2)
plot(res1, fun='cloglog', lwd=2, lty=1:2)
legend('topleft',c('Placebo','Treatment'),lty=1:2)
leukemia$cat<-rep(2,42)</pre>
leukemia$cat[leukemia$logWBC<=2.3]<-1</pre>
leukemia$cat[leukemia$logWBC>3.0]<-3</pre>
leukemia$cat<-factor(leukemia$cat,labels=c('low','median','high'))</pre>
res2<-survfit(Surv(Survt,Relapse)~cat, data=leukemia)</pre>
plot(res2, fun="cloglog", lwd=2, lty=1:3)
legend('topleft',c('low','median','high'),lty=1:3)
res3<-survfit(Surv(Survt,Relapse)~Sex,data=leukemia)
plot(res3, fun='cloglog', ylim=c(-3,1), lwd=2, lty=1:2)
legend('topleft',c('Female','Male'),lwd=2,lty=1:2)
```









Problems with log-log survival curve approach:

How parallel is parallel? Recommend:

- subjective decision
- conservative strategy: assume PH is OK unless strong evidence of nonparallelism

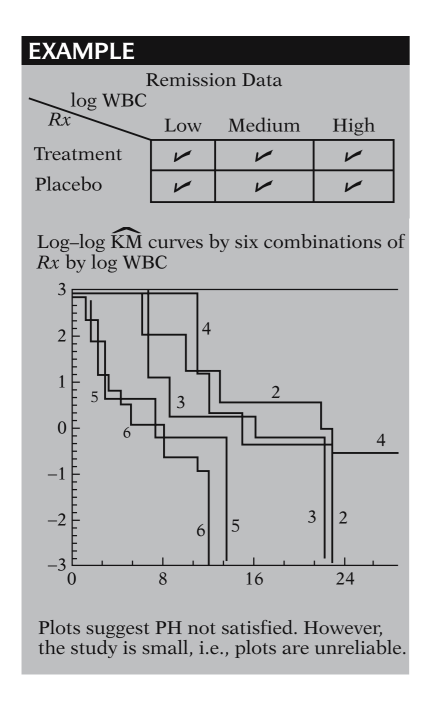
How to categorize a continuous variable?

- many categories ⇒ data "thins out"
- different categorizations may give different graphical pictures

Recommend:

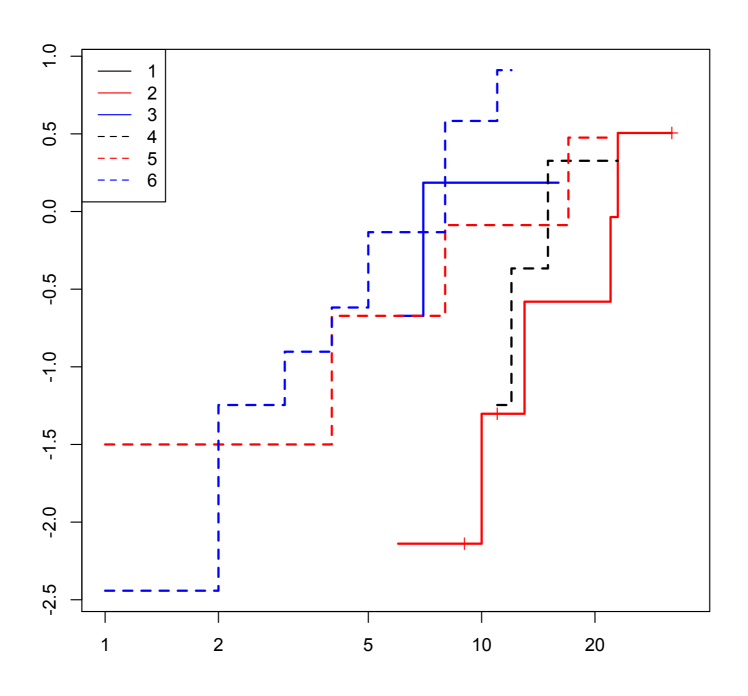
- small # of categories (2 or 3)
- meaningful choice
- reasonable balance (e.g., terciles)

- The above examples suggest that there are some problems associated with this graphical approach for assessing the PH assumption. The main problem concerns how to decide "how parallel is parallel?" This decision can be quite subjective for a given data set, particularly if the study size is relatively small. We recommend that one should use a conservative strategy for this decision by assuming the PH assumption is satisfied unless there is strong evidence of nonparallelism of the log-log curves.
- Another problem concerns how to categorize a continuous variable like log WBC. If many categories are chosen, the data "thins out" in each category, making it difficult to compare different curves. [Also, one categorization into, say, three groups may give a different graphical picture from a different categorization into three groups.]
- In categorizing continuous variables, we recommend that the number of categories be kept reasonably small (e.g., two or three) if possible, and that the choice of categories be as meaningful as possible and also provide reasonable balance of numbers (e.g., as when using terciles).



- As an example of this strategy, suppose we use the remission data again and consider both Rx and log WBC together. Because we previously had two categories of Rx and three categories of log WBC, we get a total of six combined categories, consisting of treated subjects with low log WBC, placebo subjects with low log WBC, treated subjects with medium log WBC, and so on.
- The computer results are shown here for the log-log curves corresponding to each of the six combinations of Rx with log WBC. Notice that there are several points of intersection among the six curves. Therefore, these results suggest that the PH assumption is not satisfied when considering Rx and log WBC together.
- However, the sample sizes used to estimate these curves are quite small, ranging between four subjects for group 4 (Rx=1,log WBC = low) to twelve subjects for group 6 (Rx = 1, log WBC = high), with the total study size being 42. Thus, for this small study, the use of six log-log curves provides unreliable information for assessing the PH assumption.

```
library(survival) # loading survival package
leukemia<-read.csv('leukemia.csv',header=T)</pre>
attach(leukemia)
cat < -rep(2,42)
cat[logWBC<=2.3]<-1
cat[logWBC>3.0]<-3
leukemia$cat<-factor(cat,labels=c('low','median','high'))</pre>
comb < -rep(0,42)
comb[Rx==0 & cat==1]<-1
comb[Rx==0 & cat==2]<-2
comb[Rx==0 & cat==3]<-3
comb[Rx==1 \& cat==1]<-4
comb[Rx==1 & cat==2]<-5
comb[Rx==1 & cat==3]<-6
leukemia$comb<-comb</pre>
res<-survfit(Surv(Survt,Relapse)~comb)</pre>
plot(res, fun='cloglog', lwd=2)
plot(res, fun='cloglog', lwd=2, lty=rep(1:2, each=3), col=c('black', 'red', 'blue', 'black', 'red', 'blue'))
legend('topleft',paste(1:6),lty=rep(1:3,each=3),col=c('black','red','blue','black','red','blue'))
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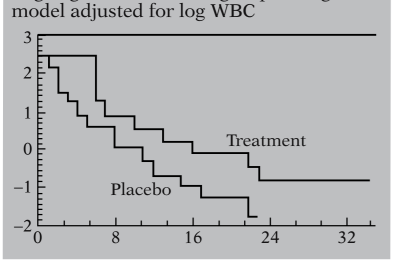
Alternative strategy: Adjust for predictors already satisfying PH assumption, i.e., use adjusted log—log Ŝ curves An alternative graphical strategy for considering several predictors together is to assess the PH assumption for one predictor adjusted for other predictors that are assumed to satisfy the PH assumption. Rather than using Kaplan–Meier curves, this involves a comparison of adjusted log–log survival curves.

EXAMPLE

Remission data:

- compare *Rx* categories adjusted for log WBC
- fit PH model for each Rx stratum
- obtain adjusted survival curves using overall mean of log WBC

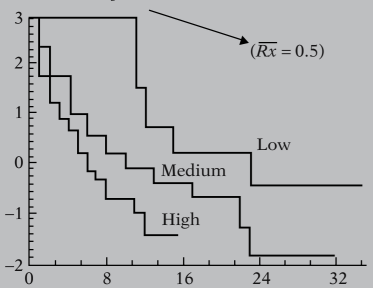
Log-log \hat{S} curves for Rx groups using PH model adjusted for log WBC



- As an example, again we consider the remission data and the predictors Rx and log WBC. To assess the PH assumption for Rx adjusted for log WBC, we would compare adjusted log-log survival curves for the two treatment categories, where each adjusted curve is derived from a PH model containing log WBC as a predictor. In computing the adjusted survival curve, we need to stratify the data by treatment, fit a PH model in each stratum, and then obtain adjusted survival probabilities using the overall mean log WBC in the estimated survival curve formula for each stratum.
- For the remission data example, the estimated log-log survival curves for the two treatment groups adjusted for log WBC are shown here. Notice that these two curves are roughly parallel, indicating that the PH assumption is satisfied for treatment.

EXAMPLE (continued)

Log-log \hat{S} curves for log WBC groups using PH model adjusted for Rx

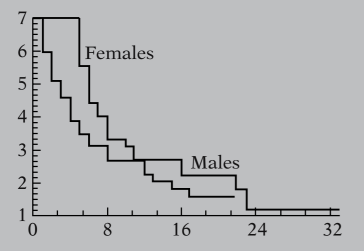


Remission data:

Assess PH assumption for Sex:

- use PH model containing *Rx* and log WBC
- use *Rx* and log WBC in survival probability formula

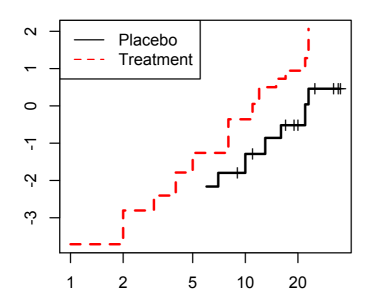
Log-log \hat{S} curves for Sex adjusted for Rx and log WBC

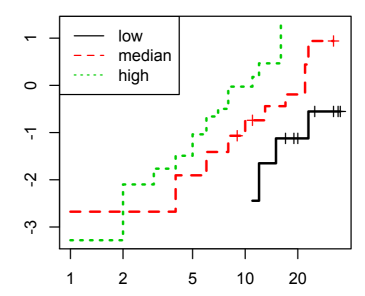


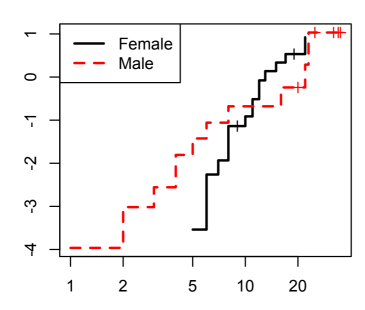
- As another example, we consider adjusted log-log survival curves for three categories of log WBC, adjusted for the treatment status (Rx) variable. The adjusted survival probabilities in this case use the overall mean Rx score, i.e., 0.5, the proportion of the 42 total subjects that are in the placebo group (i.e., half the subjects have a score of Rx = 1).
- The three log-log curves adjusted for treatment status are shown here. Although two of these curves intersect early in follow-up, they do not suggest a strong departure from parallelism overall, suggesting that the PH assumption is reasonable for log WBC, after adjusting for treatment status.
- As a third example, again using the remission data, we assess the PH assumption for Sex, adjusting for both treatment status and log WBC in the model. This involves obtaining log-log survival curves for males and females separately, using a PH model that contains both treatment status and log WBC. The adjustment uses the overall mean treatment score and the overall mean log WBC score in the formula for the estimated survival probability.
- The estimated log-log survival curves for Sex, adjusted for treatment and log WBC are shown here. These curves clearly cross, indicating that the PH assumption is not satisfied for Sex, after adjusting for treatment and log WBC.

23

```
library(survival) # loading survival package
leukemia<-read.csv('leukemia.csv',header=T)</pre>
attach(leukemia)
par(mfrow=c(2,2))
res1<-coxph(Surv(Survt,Relapse)~strata(Rx)+logWBC)
plot(survfit(res1), fun='cloglog', lwd=2, lty=1:2, col=1:2)
legend('topleft',c('Placebo','Treatment'),lty=1:2, col=1:2)
cat < -rep(2,42)
cat[logWBC<=2.3]<-1
cat[logWBC>3.0]<-3
leukemia$cat<-factor(cat,labels=c('low','median','high'))</pre>
res2<-coxph(Surv(Survt,Relapse)~strata(cat)+Rx)</pre>
plot(survfit(res2), fun='cloglog', lwd=2, lty=1:3, ,col=1:3)
legend('topleft',c('low','median','high'),lty=1:3,col=1:3)
res3<-coxph(Surv(Survt,Relapse)~Rx+logWBC+strata(Sex))
plot(survfit(res3), fun='cloglog', lwd=2, lty=1:2, col=1:2)
legend('topleft',c('Female','Male'),lwd=2,lty=1:2,col=1:2)
```







- ✓ 1. log-log survival curves
 - 2. observed versus expected survival curves

 We have thus described and illustrated one of the two graphical approaches for checking the PH assumption, that is, using log-log survival plots. In the next section, we describe an alternative approach that compares "observed" with "expected" survival curves.

Graphical analog of GOF test

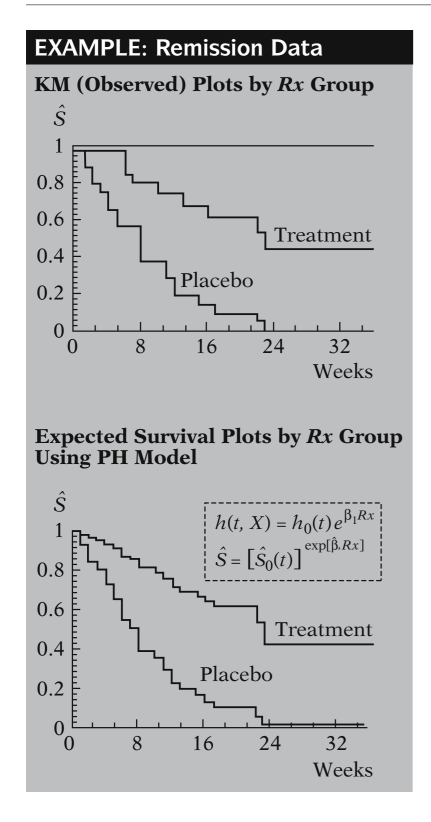
Two strategies:

- 1. One-at-a-time: uses KM curves to obtain observed plots
- 2. Adjusting for other variables: uses stratified Cox PH model to obtain observed plots (see Chapter 5)

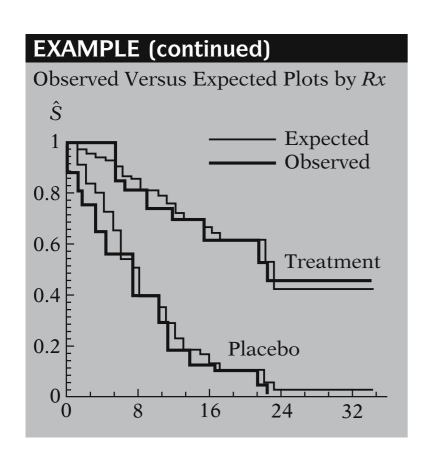
One-at-a-time:

- stratify data by categories of predictor
- obtain KM curves for each category

- The use of observed versus expected plots to assess the PH assumption is the graphical analog of the goodness-of-fit (GOF) testing approach to be described later, and is therefore a reasonable alternative to the log-log survival curve approach.
- As with the log-log approach, the observed versus expected approach may be carried out using either or both of two strategies— (1) assessing the PH assumption for variables one-at-a-time, or (2) assessing the PH assumption after adjusting for other variables. The strategy which adjusts for other variables uses a stratified Cox PH model to form observed plots, where the PH model contains the variables to be adjusted and the stratified variable is the predictor being assessed. The stratified Cox procedure is described in Chapter 5.
- Here, we describe only the one-at-a-time strategy, which involves using KM curves to obtain observed plots.
- Using the one-at-a-time strategy, we first must stratify our data by categories of the predictor to be assessed. We then obtain observed plots by deriving the KM curves separately for each category.



- As an example, for the remission data on 42 leukemia patients we have illustrated earlier, the KM plots for the treatment and placebo groups, with 21 subjects in each group, are shown here. These are the "observed" plots.
- To obtain "expected" plots, we fit a Cox PH model containing the predictor being assessed. We obtain expected plots by separately substituting the value for each category of the predictor into the formula for the estimated survival curve, thereby obtaining a separate estimated survival curve for each category.
- As an example, again using the remission data, we fit the Cox PH model with Rx as its only variable. Using the corresponding survival curve formula for this Cox model, as given in the box at the left, we then obtain separate expected plots by substituting the values of 0 (for treatment group) and 1 (for placebo group). The expected plots are shown here.



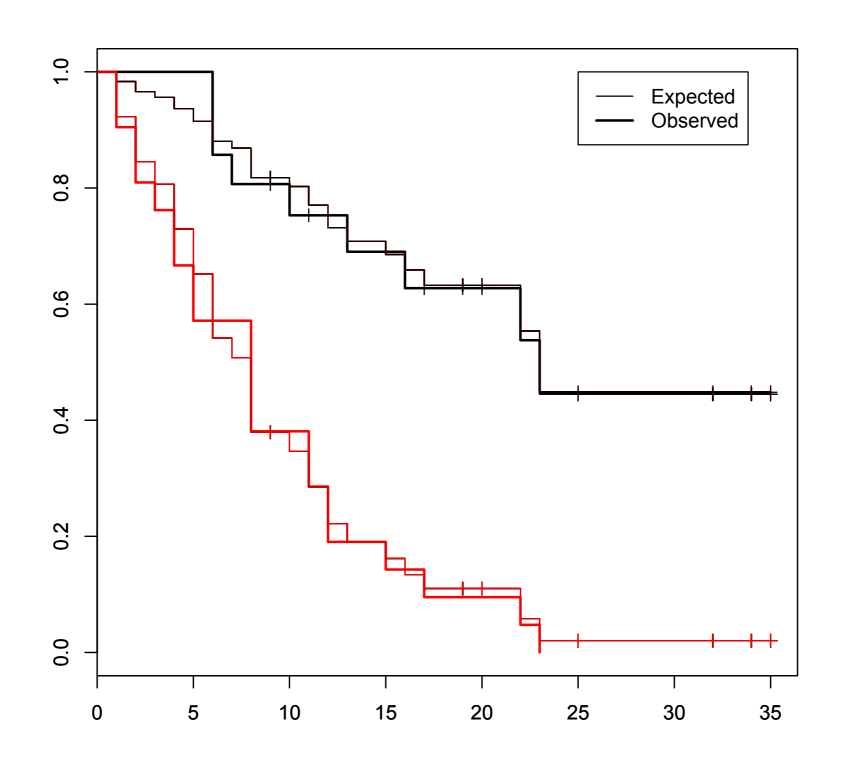
 To compare observed with expected plots we then put both sets of plots on the same graph as shown here.

If observed and expected plots are:

- **close**, complies with PH assumption
- **discrepant**, PH assumption violated

 If for each category of the predictor being assessed, the observed and expected plots are "close" to one another, we then can conclude that the PH assumption is satisfied. If, however, one or more categories show quite discrepant observed and expected plots, we conclude that the PH assumption is violated.

```
library(survival) # loading survival package
leukemia<-read.csv('leukemia.csv',header=T)</pre>
attach(leukemia)
res.km<-survfit(Surv(Survt,Relapse)~Rx,data=leukemia)</pre>
res.cox<-survfit(coxph(Surv(Survt,Relapse)~Rx,data=leukemia),newdata=leukemia)
plot(res.km, col=1:2, lwd=2)
par(new=TRUE)
plot(res.cox,col=1:2,lwd=1,main='Observed Versus Expected Plots by Rx',ylab=expression(hat(S)),xlab='')
legend(25,1,c('Expected','Observed'),lwd=1:2)
```



EXAMPLE: Remission Data (continued)

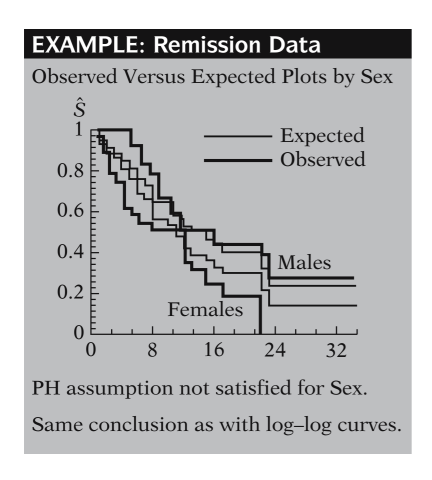
Observed and expected plots are close for each treatment group.

Conclude PH assumption not violated.

Drawback: How close is close?

Recommend: PH not satisfied *only* when plots are strongly discrepant.

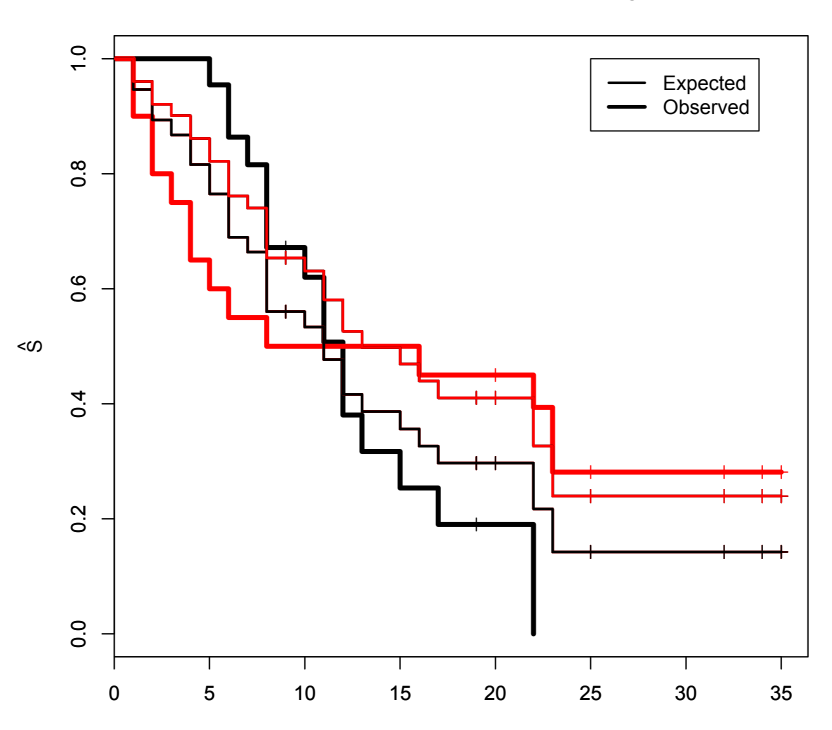
- For the example shown above, observed and expected curves appear to be quite close for each treatment group. Thus, we would conclude using this graphical approach that the treatment variable satisfies the PH assumption.
- An obvious drawback to this graphical approach is deciding "how close is close" when comparing observed versus expected curves for a given category. This is analogous to deciding "how parallel is parallel" when comparing log-log survival curves. Here, we recommend that the PH assumption be considered as not satisfied only when observed and expected plots are strongly discrepant.



- As another example, again using the remission data, we consider observed versus expected plots by Sex, as shown here. Note that the observed plots for males and females, which are described by the thicker lines, cross at about 12 weeks, whereas the expected plots don't actually intersect, with the female plot lying below the male plot throughout follow-up. Moreover, for males and females separately, the observed and expected plots are quite different from one another.
- Thus, the above plots suggest that the PH assumption is not satisfied for the variable Sex. We came to the same conclusion when using log-log survival curves, which crossed one another and were therefore clearly nonparallel.

```
library(survival) # loading survival package
leukemia<-read.csv('leukemia.csv',header=T)</pre>
attach(leukemia)
res.km<-survfit(Surv(Survt,Relapse)~Sex,data=leukemia)</pre>
res.cox<-survfit(coxph(Survt,Relapse)~Sex,data=leukemia),newdata=leukemia)
plot(res.km, col=1:2, lwd=4)
par(new=TRUE)
plot(res.cox,col=1:2,lwd=2,main='Observed Versus Expected Plots by Sex',ylab=expression(hat(S)),xlab='')
legend(25,1,c('Expected','Observed'),lwd=2:4)
```

Observed Versus Expected Plots by Sex



Continuous variable:

- form strata from categories
- observed plots are KM curves for each category
- two options for expected plots
 - 1. Use PH model with k-1 dummy variables X_i for k categories, i.e.,

$$h(t, \mathbf{X}) = h_0(t) \exp\left(\sum_{i=1}^{k-1} \beta_i X_i\right)$$

Obtain adjusted survival curve:

$$\hat{S}(t, \mathbf{X}_c) = [\hat{S}_0(t)]^{\exp(\sum \hat{\beta}_i X_{ci})}$$

where

$$\mathbf{X}_c = (X_{c1}, X_{c2}, \dots, X_{c,k-1})$$
 gives values of dummy variables for category c .

- When using observed versus expected plots to assess the PH assumption for a continuous variable, observed plots are derived, as for categorical variables, by forming strata from categories of the continuous variable and then obtaining KM curves for each category.
- However, for continuous predictors, there are two options available for computing expected plots. One option is to use a Cox PH model which contains k – 1 dummy variables to indicate k categories. The expected plot for a given category is then obtained as an adjusted survival curve by substituting the values for the dummy variables that define the given category into the formula for the estimated survival curve, as shown here for category c.

Options for a continuous variable:

2. Use PH model:

$$h(t,X) = h_0(t) \exp(\beta X)$$
Continuous

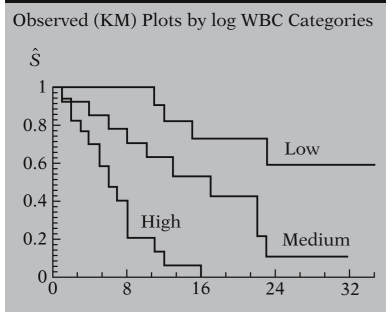
Obtain adjusted survival curve:

$$\hat{S}(t, \overline{X}_c) = [\hat{S}_0(t)]^{\exp(\hat{\beta}\,\overline{X}_c)}$$

where \overline{X}_c denotes the mean value for the variable X within category c.

 The second option is to use a Cox PH model containing the continuous predictor being assessed. Expected plots are then obtained as adjusted survival curves by specifying predictor values that distinguish categories, as, for example, when using mean predictor values for each category.

EXAMPLE: Remission Data



Option 1:

$$h(t, \mathbf{X}) = h_0(t) \exp(\beta_1 X_1 + \beta_2 X_2)$$
where $X_1 = \begin{cases} 1 & \text{if high} \\ 0 & \text{if other} \end{cases}$ $X_2 = \begin{cases} 1 & \text{if medium} \\ 0 & \text{if other} \end{cases}$

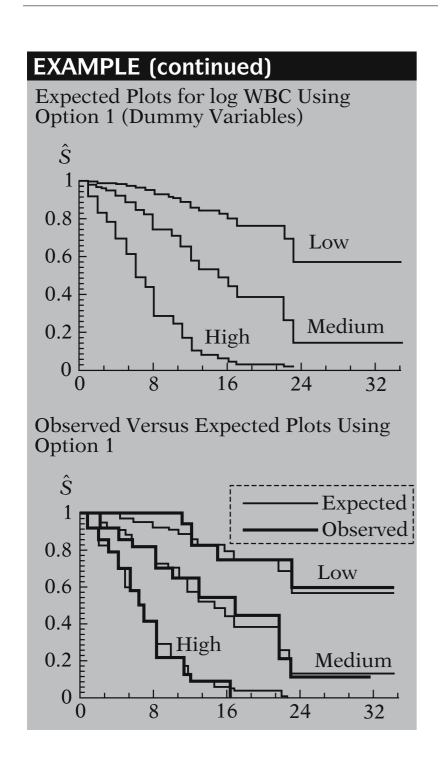
so that

high =
$$(1, 0)$$
; medium = $(0, 1)$; low = $(0, 0)$

Expected survival plots:

$$X_1 = 1, X_2 = 0$$
: $\hat{S}(t, X_{\text{high}}) = [\hat{S}_0(t)]^{\exp(\hat{\beta}_1)}$
 $X_1 = 0, X_2 = 1$: $\hat{S}(t, X_{\text{medium}}) = [\hat{S}_0(t)]^{\exp(\hat{\beta}_2)}$
 $X_1 = 0, X_2 = 0$: $\hat{S}(t, X_{\text{low}}) = [\hat{S}_0(t)]$

- As an example to illustrate both options, we consider the continuous variable log WBC from the remission data example. To assess the PH assumption for this variable, we would first stratify log WBC into, say, three categories—low, medium, and high. The observed plots would then be obtained as KM curves for each of the three strata, as shown here.
- Using option 1, expected plots would be obtained by fitting a Cox PH model containing two dummy variables X_1 and X_2 , as shown here, where X_1 takes the values 1 if high or 0 if other and X_2 takes the values 1 if medium or 0 if other. Thus, when log WBC is high, the values of X_1 and X_2 are 1 and 0, respectively; whereas when log WBC is medium, the values are 0 and 1, respectively; and when log WBC is low, the values are both 0.
- The expected survival plots for high, medium, and low categories are then obtained by substituting each of the three specifications of X_1 and X_2 into the formula for the estimated survival curve, and then plotting the three curves.



 The expected plots using option 1 (the dummy variable approach) are shown here for the three categories of log WBC.

Here we put the observed and expected plots on the same graph.
 Although there are some discrepancies, particularly early in follow-up for the low log WBC category, these plots suggest overall that the PH assumption is satisfied for log WBC.

EXAMPLE (continued)

Option 2: Treat log WBC as continuous $h(t, \mathbf{X}) = h_0(t) \exp[\beta(\log \text{WBC})]$

$$\log \text{WBC}_{\text{high}} = 3.83$$
:
 $\hat{S}(t, X_{\text{high}}) = [\hat{S}_0(t)] \exp[3.83\hat{\beta}]$

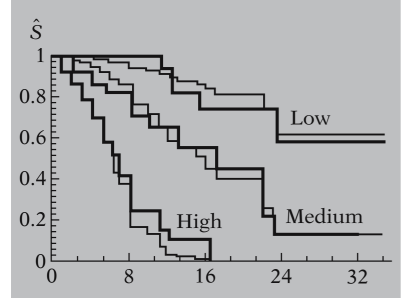
$$\overline{\log \text{WBC}}_{\text{med}} = 2.64$$
:

$$\hat{S}(t, X_{\text{med}}) = [\hat{S}_0(t)] \exp[2.64\hat{\beta}]$$

$$\overline{\log \text{WBC}}_{\text{low}} = 1.71$$
:

$$\hat{S}(t, X_{\text{low}}) = [\hat{S}_0(t)]^{\exp[1.71\hat{\beta}]}$$

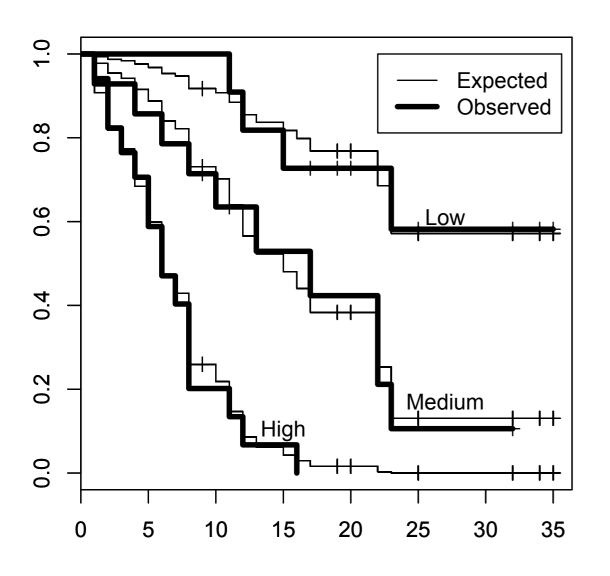
Observed Versus Expected Plots for log WBC Using Option 2



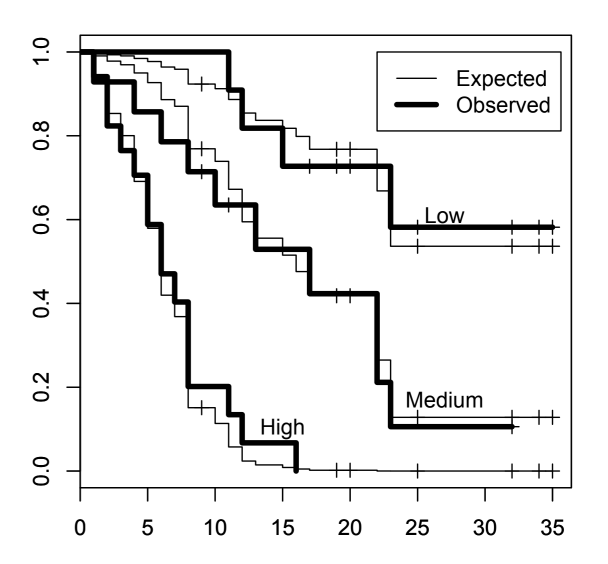
- Using option 2, expected plots would be obtained by first fitting a Cox PH model containing the continuous variable log WBC, as shown here.
- Adjusted survival curves are then obtained for specified values of log WBC that summarize the three categories used to form observed curves. Here, we find that the mean log WBC scores for low, medium, and high categories are, respectively, 1.71, 2.64, and 3.83. These values are substituted into the estimated survival curve formula as shown here.
- Here are the observed and expected plots using option 2. As with option 1, although there are some discrepancies within categories, overall, these plots suggest that the PH assumption is satisfied for the log WBC variable.

```
library(survival) # loading survival package
leukemia<-read.csv('leukemia.csv',header=T)</pre>
attach(leukemia)
cat<-rep('medium',42); cat[logWBC<=2.3]<-'low'; cat[logWBC>3.0]<-'high'</pre>
leukemia$cat<-cat
res.km<-survfit(Surv(Survt,Relapse)~cat,data=leukemia)
# option 1
res.cox<-survfit(coxph(Survt,Relapse)~cat,data=leukemia),newdata=leukemia)
par(mfrow=c(1,2))
plot(res.km, lwd=4)
par(new=TRUE)
plot(res.cox, lwd=1, main='Observed Versus Expected Plots by Option 1')
legend(22,1,c('Expected','Observed'),lwd=c(1,4))
text(27,0.61,'Low')
text(27,0.17, 'Medium')
text(15,0.1, 'High')
# option 2
m1<-mean(logWBC[cat=='low']); m2<-mean(logWBC[cat=='medium']); m3<-mean(logWBC[cat=='high'])</pre>
leukemia2<-data.frame(logWBC=c(m1,m2,m3))</pre>
res.cox<-survfit(coxph(Survt,Relapse)~logWBC,data=leukemia),newdata=leukemia2)
par(new=FALSE)
plot(res.km, lwd=4)
par(new=TRUE)
plot(res.cox,lwd=1,main='Observed Versus Expected Plots by Option 2')
legend(22,1,c('Expected','Observed'),lwd=c(1,4))
text(27,0.61, 'Low')
text(27,0.17, 'Medium')
text(15,0.1, 'High')
```

Observed Versus Expected Plots by Option 1



Observed Versus Expected Plots by Option 2



Statistical test appealing

- Provides p-value
- More objective decision than when using graphical approach

Test of Grambsch and Therneau (1994)

- Variation of test of Schoenfeld
- Uses Schoenfeld residuals

- The GOF testing approach is appealing because it provides a test statistic and p-value for assessing the PH assumption for a given predictor of interest. Thus, the researcher can make a more objective decision using a statistical test than is typically possible when using either of the two graphical approaches described above.
- A number of different tests for assessing the PH assumption have been proposed in the literature. We present the test of Grambsch and Therneau (1994), a variation of a test originally proposed by Schoenfeld (1982) and based on the residuals defined by Schoenfeld, now called the Schoenfeld residuals.

Schoenfeld residuals defined for

- Each predictor in model
- Every subject who has event

Consider Cox PH model

$$h(t) = h_0(t) \exp(\beta_1 RX + \beta_2 \log WBC + \beta_3 SEX)$$

3 predictors → 3 Schoenfeld residuals for each subject who has event

Schoenfeld residual for ith subject for LOGWBC

- Observed LOGWBC
- LOGWBC weighted average

Weights are other subjects' hazard (from subjects still at risk)

Underlying idea of test If PH holds then Schoenfeld residuals uncorrelated with time

- For each predictor in the model, Schoenfeld residuals are defined for every subject who has an event. For example, consider a Cox PH model with three predictors: RX, LOGWBC, and SEX. Then there are three Schoenfeld residuals defined for each subject who has an event, one for each of the three predictors.
- Suppose subject i has an event at time t_j . Then her Schoenfeld residual for LOGWBC is her observed value of log white blood cell count minus a weighted average of the log white blood cell counts for the other subjects still at risk at time t_j . The weights are each subject's hazard.
- The idea behind the statistical test is that if the PH assumption holds for a particular covariate then the Schoenfeld residuals for that covariate will not be related to survival time.

Steps for test implementation

- 1. Obtain Schoenfeld residuals
- 2. Rank failure times
- 3. Test correlation of residuals to ranked failure time H_0 : $\rho = 0$

H₀ rejected Conclude PH assumption violated

R uses scaled Schoenfeld residuals rather than Schoenfeld residuals (typically similar results) The implementation of the test can be thought of as a three-step process.

Step 1. Run a Cox PH model and obtain Schoenfeld residuals for each predictor.

Step 2. Create a variable that ranks the order of failures. The subject who has the first (earliest) event gets a value of 1, the next gets a value of 2, and so on.

Step 3. Test the correlation between the variables created in the first and second steps. The null hypothesis is that the correlation between the Schoenfeld residuals and ranked failure time is zero.

- Rejection of the null hypothesis leads to a conclusion that the PH assumption is violated.
- The implementation of the test for the PH assumption in R is done by cox.zph() function. R uses a slight variation of the test we just described in that it uses the scaled Schoenfeld residual rather than the Schoenfeld residual (Grambsch and Therneau, 1994). The tests typically (but not always) yield similar results.

EXAMPLE: Remisson Data

Both variables satisfy PH assumption.

Note: p=0.981 assesses PH for Rx, assuming PH OK for log WBC.

```
> cox.zph(res2)

rho chisq p

Rx -0.1017 0.344 0.5578

logWBC 0.0595 0.161 0.6883

Sex -0.3684 4.076 0.0435

GLOBAL NA 4.232 0.2374
```

log WBC and Rx satisfy PH.

Sex does not satisfy PH.

(Same conclusion using graphical approaches).

- To illustrate the statistical test approach, we return to the remission data example. The printout on the left gives p-values for treatment group and log WBC variables based on fitting a Cox PH model containing these two variables.
- The p-values are quite high for both variables, suggesting that both variables satisfy the PH assumption. Note that each of these p-values tests the assumption for one variable given that the other predictors are included in the model. For example, the p-value of 0.981 assesses the PH assumption for Rx, assuming the PH assumption is satisfied for log WBC.
- As another example, consider the computer results shown here for a Cox PH model containing the variable SEX in addition to log WBC and treatment group. The p-values for log WBC and treatment group are still nonsignificant. However, the p-value for SEX is significant below the 0.05 level, while the global test (on 3 degrees of freedom) is not quite statistically significant. This result suggests that log WBC and treatment group satisfy the PH assumption, whereas SEX does not. We came to the same conclusion about these variables using the graphical procedures described earlier.

```
library(survival) # loading survival package
leukemia<-read.csv('leukemia.csv',header=T)</pre>
attach(leukemia)
res1<-coxph(Surv(Survt,Relapse)~Rx+logWBC,data=leukemia,x=TRUE)
cox.zph(res1)
res2<-coxph(Surv(Survt,Relapse)~Rx+logWBC+Sex,data=leukemia,x=TRUE)
cox.zph(res2)
```

Statistical Tests

Null is never proven

May say not enough evidence to reject

p-value can be driven by sample size

- Small sample—gross violation of null may not be significant
- Large sample—slight violation of null may be highly significant

Test—more objective
Graph—more objective, but can
detect specific violations

Recommend—Use both graphs and tests

• An important point concerning a testing approach is that the null hypothesis is never proven with a statistical test. The most that may be said is that there is not enough evidence to reject the null. A pvalue can be driven by sample size. A gross violation of the null assumption may not be statistically significant if the sample is very small. Conversely, a slight violation of the null assumption may be highly significant if the sample is very large.

• A statistical test offers a more objective approach for assessing the PH assumption compared to the subjectivity of the graphical approach. However, the graphical approach enables the researcher to detect specific kinds of departures from the PH assumption; the researcher can see what is going on from the graph. Consequently, we recommend that when assessing the PH assumption, the investigator use both graphical procedures and statistical testing before making a final decision.

Extended Cox model: contains product terms of the form $X \times g(t)$, where g(t) is a function of time.

One-at-a-time model:

$$h(t, \mathbf{X}) = h_0(t) \exp[\beta X + \delta X \times g(t)]$$

Some choices for g(t):

$$g(t) = t$$

$$g(t) = \log t$$

$$g(t) = \begin{cases} 1 & \text{if } t \ge t_0 \\ 0 & \text{if } t < t_0 \end{cases} \text{ (heaviside function)}$$

 H_0 : $\delta = 0$ Under H_0 , the model reduces to:

$$h(t, \mathbf{X}) = h_0(t) \exp[\beta X]$$

Use either Wald statistic or likelihood ratio statistic: \mathcal{X}^2 with 1 df under H_0

- When time-dependent variables are used to assess the PH assumption for a time-independent variable, the Cox model is extended to contain product (i.e., interaction) terms involving the time-independent variable being assessed and some function of time.
- When assessing predictors one-at-a-time, the extended Cox model takes the general form shown here for the predictor *X*.
- One choice for the function g(t) is simply g(t) equal to t, so that the product term takes the form $X \times t$. Other choices for g(t) are also possible, for example, $\log t$.
- Using the above one-at-a-time model, we assess the PH assumption by testing for the significance of the product term. The null hypothesis is therefore " δ equal to zero." Note that if the null hypothesis is true, the model reduces to a Cox PH model containing the single variable X.
- The test can be carried out using either a Wald statistic or a likelihood ratio statistic. In either case, the test statistic has a chisquare distribution with one degree of freedom under the null hypothesis.

EXAMPLE

 $h(t, \mathbf{X}) = h_0(t) \exp[\beta_1 \operatorname{Sex} + \beta_2 (\operatorname{Sex} \times t)]$

 $\beta_2 \neq 0 \Rightarrow PH$ assumption violated

Strategies for assessing PH:

- one-at-a-time
- several predictors simultaneously
- for a given predictor adjusted for other predictors

- For example, if the PH assumption is being assessed for Sex, a Cox model might be extended to include the variable Sex × t in addition to Sex. If the coefficient of the product term turns out to be significant, we can conclude that the PH assumption is violated for Sex.
- In addition to a one-at-a-time strategy, the extended Cox model can also be used to assess the PH assumption for several predictors simultaneously as well as for a given predictor adjusted for other predictors in the model.

Several predictors simultaneously:

$$h(t, \mathbf{X}) = h_0(t) \exp\left(\sum_{i=1}^p [\beta_i X_i + \delta_i X_i \times g_i(t)]\right)$$

 $g_i(t)$ = function of time for ith predictor

$$H_0$$
: $\delta_1 = \delta_2 = \dots = \delta_p = 0$

$$LR = -2 \ln L_{\text{PH model}}$$

$$-(-2 \ln L_{\text{ext. Cox model}})$$

$$\sim \mathcal{X}_p^2 \text{ under } H_0$$

Cox PH (reduced) model:

$$h(t, \mathbf{X}) = h_0(t) \exp\left(\sum_{i=1}^p \beta_i X_i\right)$$

• To assess the PH assumption for several predictors simultaneously, the form of the extended model is shown here. This model contains the predictors being assessed as main effect terms and also as product terms with some function of time. Note that different predictors may require different functions of time; hence, the notation g_i(t) is used to define the time function for the ith predictor.

• With the above model, we test for the PH assumption simultaneously by assessing the null hypothesis that all the δ_i coefficients are equal to zero. This requires a likelihood ratio chisquare statistic with p degrees of freedom, where p denotes the number of predictors being assessed. The LR statistic computes the difference between the log likelihood statistic—-2 ln L—for the PH model and the log likelihood statistic for the extended Cox model. Note that under the null hypothesis, the model reduces to the Cox PH model shown here.

EXAMPLE: Remission Data

$$h(t, \mathbf{X}) = h_0(t) \exp \left[\beta_1 (Rx) + \beta_2 (\log \text{WBC}) + \beta_3 (\text{Sex}) + \delta_1 (Rx) \times g(t) + \delta_2 (\log \text{WBC}) \times g(t) + \delta_3 (\text{Sex}) \times g(t)\right]$$
where $g(t) = \begin{cases} 1 & \text{if } t \ge 7 \\ 0 & \text{if } t < 7 \end{cases}$

$$H_0: \delta_1 = \delta_2 = \delta_3 = 0$$

$$LR \stackrel{\sim}{\sim} \chi^2 \text{ with } 3 \text{ df}$$

If test is significant, use backward elimination to find predictors not satisfying PH assumption.

- As an example, we assess the PH assumption for the predictors Rx, log WBC, and Sex from the remission data considered previously. The extended Cox model is given as shown here, where the functions $g_i(t)$ have been chosen to be the same "heaviside" function defined by g(t) equals 1 if t is 7 weeks or more and g(t) equals 0 if t is less than 7 weeks. The null hypothesis is that all three δ coefficients are equals to zero. The test statistic is a likelihoodratio chi-square with 3 degrees of freedom.
- If the above test is found to be significant, then we can conclude that the PH assumption is not satisfied for at least one of the predictors in the model. To determine which predictor(s) do not satisfy the PH assumption, we could proceed by backward elimination of nonsignificant product terms until a final model is attained.

Heavyside function:

$$g(t) = \begin{cases} 1 & \text{if } t \ge 7 \\ 0 & \text{if } t < 7 \end{cases}$$

 $h(t, \mathbf{X})$ differs for $t \ge 7$ and t < 7.

Properties of heaviside functions and numerical results are described in Chapter 6.

Assessing PH for a given predictor adjusted for other predictors:

$$h(t, \mathbf{X}) = h_0(t) \exp \left[\sum_{i=1}^{p-1} \beta_i X_i + \beta^* X^* \right]$$

$$+\delta^*X^*\times g(t)$$

 $X^* = \text{Predictor of interest}$ $H_0: \delta^* = 0$ Wald or LR statistic $\sim \chi^2$ with 1 df • Note that the use of a heaviside function for g(t) in the above example yields different expressions for the hazard function depending on whether t is greater than or equal to 7 weeks or t is less than 7 weeks. Chapter 6 provides further details on the properties of heaviside functions, and also provides numerical results from fitting extended Cox models.

• We show here an extended Cox model that can be used to evaluate the PH assumption for a given predictor adjusted for predictors already satisfying the PH assumption. The predictor of interest is denoted as X^* , and the predictors considered to satisfy the PH assumption are denoted as X_i . The null hypothesis is that the coefficient δ^* of the product term $X^*g(t)$ is equal to zero. The test statistic can either be a Wald statistic or a likelihood ratio statistic, with either statistic having a chi-square distribution with 1 degree of freedom under the null hypothesis.

EXAMPLE: Remission Data

```
For Sex, adjusted for Rx and log WBC:

h(t, \mathbf{X}) = h_0(t) \exp[\beta_1 (Rx) + \beta_2 (\log \text{WBC}) + \beta^* (\text{Sex}) + \delta^* (\text{Sex}) \times g(t)
```

Two models for LR test of PH:

- 1. Cox PH model
- 2. extended Cox model

Drawback: choice of $g_i(t)$

Different choices may lead to different conclusions about PH assumption.

- As an example, suppose, again considering the remission data, we assess the PH assumption for the variable, Sex, adjusted for the variables Rx and log WBC, which we assume already satisfy the PH assumption. Then, the extended Cox model for this situation is shown here.
- To carry out the computations for any of the likelihood ratio tests described above, two different types of models, a PH model and an extended Cox model, need to be fit. See the Computer Appendix for details on how the extended Cox model is fit using R.
- The primary drawback of the use of an extended Cox model for assessing the PH assumption concerns the choice of the functions g_i(t) for the time-dependent product terms in the model. This choice is typically not clear-cut, and it is possible that different choices, such as g(t) equal to t versus log t versus a heaviside function, may result in different conclusions about whether the PH assumption is satisfied.

Chapter 6: Time-dependent covariates

This presentation: Three methods for assessing PH.

- i. graphical
- ii. GOF
- iii. time-dependent covariates

Recommend using at least two methods.

- Further discussion of the use of time-dependent covariates in an extended Cox model is provided in Chapter 6.
- This presentation is now complete. We have described and illustrated three methods for assessing the PH assumption: graphical, goodness-of-fit (GOF), and time-dependent covariate methods. Each of these methods has both advantages and drawbacks. We recommend that the researcher use at least two of these approaches when assessing the PH assumption.

Chapters

- 1. Introduction to Survival Analysis
- 2. Kaplan–Meier Survival Curves and the Log–Rank Test
- 3. The Cox Proportional Hazards Model and Its Characteristics
- ✓ 4. Evaluating the Proportional Hazards Assumption

Next:

- 5. The Stratified Cox Procedure
- 6. Extension of the Cox Proportional Hazards Model for Time-Dependent Variables

- We suggest that the reader review this presentation using the detailed outline that follows. Then answer the practice exercises and the test that follow.
- The next Chapter (5) is entitled "The Stratified Cox Procedure."
 There, we describe how to use a stratification procedure to fit a PH model when one or more of the predictors do not satisfy the PH assumption.