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Session 8: Survival analysis part 3

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CUNY SPH Biostatistics 2

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Learning objectives

- 1 Check model assumptions and fit of the Cox model
 - residuals analysis
 - log-minus-log plot
- 2 Fit and interpret multivariate Cox models
 - perform tests for trend
 - predict survival for specific covariate patterns
 - predict survival for adjusted coefficients
 - 3 Explain stratified analysis
- 4 Identify situations of competing risks
- 5 Describe the application of Propensity Score analysis
- Vittinghoff sections 6.2-6.4

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Outline

- Review
- 2 Assumptions of Cox PH model
- 3 Tests for trend
- 4 Predictions for specific covariate patterns
 - 5 Stratification
- 6 Competing risks
- Propensity Score analysis to control for confounding

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Cox proportional hazards model

- Cox proportional hazard regression assesses the relationship between a right-censored, time-to-event outcome and multiple predictors:
 - categorical variables (e.g., treatment groups)
 - continuous variables

$$log(HR(x_i)) = log \frac{h(t|x_i)}{h_0(t)} = \beta_0 + \beta_1 x_{1i} + \beta_2 x_{2i} + ... + \beta_p x_{pi}$$

- $HR(x_i)$ is the hazard of patient i relative to baseline
- $h(t|x_i)$ is the time-dependent hazard function h(t) for patient i
- $h_0(t)$ is the baseline hazard function, and is the negative of the slope of the $S_0(t)$, the baseline survival function.

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Caveats and Assumptions

- Categories with no events
 - can occur when the group is small or its risk is low
 - HRs with respect to such a reference group are infinite
 - hypothesis tests and CIs are difficult / impossible to interpret
- Assumptions of Cox PH model
 - Constant hazard ratio over time (proportional hazards)
 - Linear association between log(HR) and predictors (log-linearity) / multiplicative relationship between hazard and predictors
 - Independence of survival times between individuals in the sample

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Residuals analysis

- Residuals are used to investigate the lack of fit of a model to a given subject.
- For Cox regression, there's no easy analog to the usual "observed minus predicted" residual

```
library(pensim)
set.seed(1)
mydat <- create.data(</pre>
  nvars = c(1, 1),
  nsamples = 500,
  cors = c(0, 0),
  associations = c(0.5, 0.5),
  firstonly = c(TRUE, TRUE),
  censoring = c(0, 8.5)
) $data
```

Rename variables of simulated data, and make one variable cate library(dplyr)

```
##
```

Attaching package: 'dplyr'

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Simulated data to test residuals methods

summary(mydat)

```
Var1
                     Var2
                                          time
##
                                                          cens
##
    low:323
                Min.
                       :-2.99695
                                    Min.
                                                5
                                                    Min.
                                                            :0.000
##
    high:177
                1st Qu.:-0.79008
                                    1st Qu.: 691
                                                     1st Qu.:0.000
##
                Median : -0.02126
                                    Median: 1970
                                                    Median :1.000
                                            :2529
##
                Mean
                       :-0.04594
                                    Mean
                                                    Mean
                                                            :0.526
##
                3rd Qu.: 0.68933
                                    3rd Qu.:3874
                                                     3rd Qu.:1.000
                       : 3.05574
##
                Max.
                                    Max.
                                            :8481
                                                    Max.
                                                            :1.000
```

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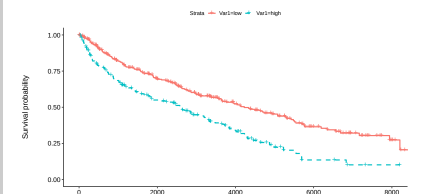
Propensity score analysis

Kaplan-Meier plot of simulated data, stratified by Var1

Loading required package: ggplot2

Loading required package: ggpubr

Warning: Vectorized input to 'element_text()' is n
Results may be unexpected or may change in future



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Martingale residuals

- censoring variable c_i (1 if event, 0 if censored) minus the estimated cumulative hazard function $H(t_i, X_i, \beta_i)$ (1 survival function)
 - E.g., for a subject censored at 1 year ($c_i = 0$), whose predicted cumulative hazard at 1 year was 30%, Martingale = 0 0.30 = -0.30.
 - E.g. for a subject who had an event at 6 months, and whose predicted cumulative hazard at 6 months was 80%, Margingale = 1 0.8 = 0.2.
- Problem: not symmetrically distributed, even when model fits the data well

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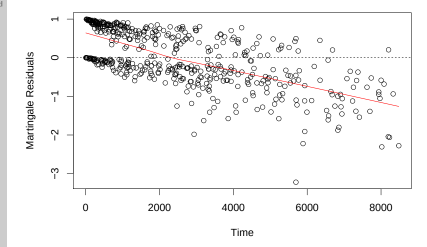
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Martingale residuals in simulated data



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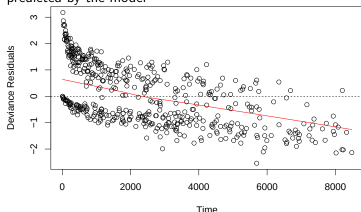
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Deviance residuals in simulated data

- Deviance residuals are scaled Martingale residuals
- Should be more symmetrically distributed about zero?
- Observations with large deviance residuals are poorly predicted by the model



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Schoenfeld residuals

- technical definition: contribution of a covariate at each event time to the partial derivative of the log-likelihood
- intuitive interpretation: the observed minus the expected values of the covariates at each event time.
- a random (unsystematic) pattern across event times gives evidence the covariate effect is not changing with respect to time
- If it is systematic, it suggests that as time passes, the covariate effect is changing.

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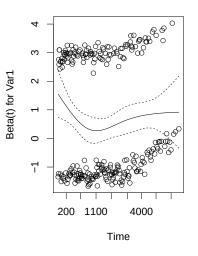
Predictions for specific covariate patterns

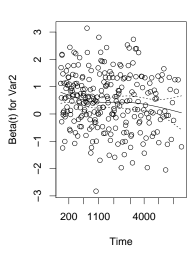
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Schoenfeld residuals for simulated data





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Schoenfeld test for proportional hazards

- Tests correlation between scaled Schoenfeld residuals and time
- Equivalent to fitting a simple linear regression model with time as the predictor and residuals as the outcome
- Parametric analog of smoothing the residuals against time using LOWESS
- If the hazard ratio is constant, correlation should be zero.
 - Positive values of the correlation suggest that the log-hazard ratio increases with time.

```
## Chisq df p
## Var1 0.00887 1 0.925
## Var2 4.92734 1 0.026
## GLOBAL 5.07415 2 0.079
```

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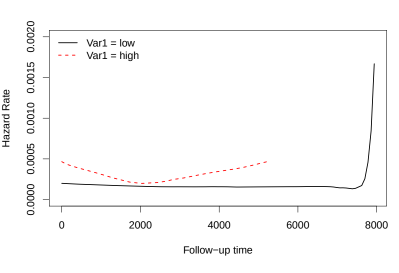
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The hazard function h(t), stratified by Var1



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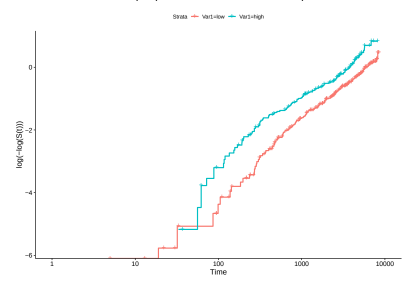
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Log-minus-log plot

• Used to check proportional hazards assumption



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Example: Primary Biliary Cirrhosis (PBC)

- Mayo Clinic trial in primary biliary cirrhosis (PBC) of the liver conducted between 1974 and 1984, n=424 patients.
- randomized placebo controlled trial of the drug D-penicillamine.
 - 312 cases from RCT, plus additional 112 not from RCT.
- Primary outcome is (censored) time to death

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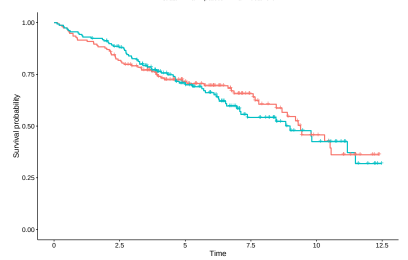
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Kaplan-Meier plot of treatment and placebo arms



Warning: Vectorized input to 'element_text()' is n

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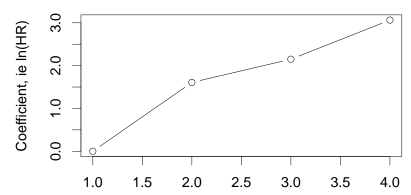
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What are tests for trend?

- For any kind of multivariate model including an ordinal variable
- Such as cancer stage (1, 2, 3, 4), age category, ...
 - Is there a linear / quadratic / cubic relationship between coefficients and their order?
 - Test by LRT or Wald Test



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How to predict survival from a Cox model?

- The Cox model is a *relative* risk model
 - only predicts relative risks between pairs of subjects
- Key is to calculate the overall S(t), then multiply it by the relative hazard for the specific covariate pattern.
- In this example we plot the baseline survival for all stages together, then for stages 1-4 separately.

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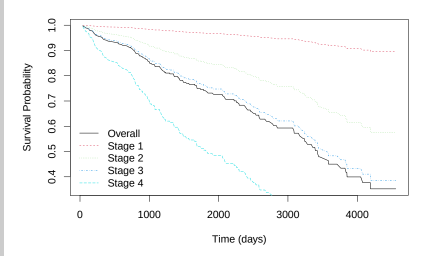
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Predicted survival for specific covariate patterns



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Multivariate regression

- Same coding and objectives as for lm() and glm()
 - controlling for confounding
 - testing for mediation
 - testing for interaction

```
Session 8:
                               + stage + arm, data = pbc.os)
                  summary(fit)
  Survival
analysis part 3
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                  ## Call:
                  ## coxph(formula = Surv(time, os) ~ age + sex + edema + stage +
                         arm, data = pbc.os)
                  ##
Learning
                  ##
objectives and
                  ##
                      n= 312, number of events= 125
outline
                 ##
                  ##
                                       coef exp(coef) se(coef)
                                                                     z Pr(>|z|)
Review
                  ## age
                                   0.027618 1.028003 0.009362 2.950 0.00318 **
                  ## sexf
                                 -0.317540 0.727938 0.248839 -1.276 0.20193
Checking
                                  0.538715 1.713804 0.275287 1.957 0.05036 .
                  ## edema0.5
assumptions
                                   2.080422
                                            8.007845 0.276959 7.512 5.84e-14 ***
                  ## edema1
of Cox model
                  ## stage2
                                  1.535263 4.642546 1.034854 1.484 0.13793
                                            7.375893 1.016097 1.967
                  ## stage3
                                  1.998217
                                                                        0.04923 *
Tests for
                  ## stage4
                                  2.666263 14.386101
                                                       1.016234 2.624
                                                                        0.00870 **
trend
                  ## armtreatment 0.057946 1.059658 0.189200 0.306 0.75940
What are
                  ## ---
tests for
                  ## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
trend?
                  ##
                 ##
                                  exp(coef) exp(-coef) lower .95 upper .95
Predictions
                                     1.0280
                                               0.97276
                                                          1.0093
                  ## age
                                                                     1.047
for specific
                  ## sexf
                                     0.7279
                                              1.37374
                                                          0.4470
                                                                     1.186
covariate
                  ## edema0.5
                                     1.7138
                                               0.58350
                                                          0.9992
                                                                     2.940
patterns
                                     8.0078
                                              0.12488
                                                          4.6534
                                                                  13.780
                  ## edema1
                                     4.6425
                                               0.21540
                                                          0.6108
                                                                    35.288
                  ## stage2
Stratification
                  ## stage3
                                    7.3759
                                               0.13558
                                                          1.0067
                                                                    54.040
                  ## stage4
                                    14.3861
                                               0.06951
                                                          1.9630
                                                                   105.430
                                     1.0597
                                               0.94370
                                                          0.7313
                                                                     1.535
                  ## armtreatment
Risks Data
                  ##
                  ## Concordance= 0.77 (se = 0.022 )
Propensity
                  ## Likelihood ratio test= 107.6 on 8 df.
                                                              p = < 2e - 16
score analysis
                  ## Wald test
                                          = 120.8 on 8 df,
                                                              p=<2e-16
                  ## Score (logrank) test = 177.1 on 8 df,
                                                              p = < 2e - 16
```

fit <- coxph(Surv(time, os) ~ age + sex + edema

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Predicted survival for adjusted coefficients

- Can create Kaplan-Meier curves for crude or unadjusted coefficients
 - Section 6.3.2.3 in Vittinghoff
- Idea is to estimate hazard ratio in an unadjusted model:

```
unadjfit <- coxph(Surv(time, os) ~ stage, data = pbc.os)
coef(unadjfit)</pre>
```

```
## stage2 stage3 stage4
## 1.607014 2.149500 3.062775
```

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Predicted survival for adjusted coefficients (cont'd)

• and in an adjusted model:

```
edema0.5
##
             age
                          sexf
                                                     edema1
                                                                   stag
##
      0.0276179
                   -0.3175396
                                   0.5387152
                                                  2.0804217
                                                                1.53526
##
          stage4 armtreatment
##
      2.6662626
                     0.0579460
```

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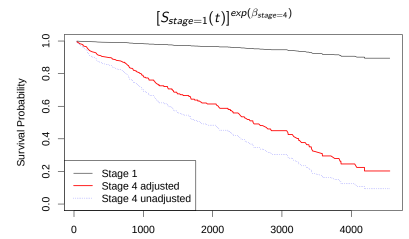
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Predicted survival for adjusted coefficients (cont'd)

 The survival function will be calculated for a "baseline" group, say stage 1, then exponentiated with the adjusted coefficient, e.g.:



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What is stratification?

- relevant to all kinds of regression, not just survival analysis
- when analysis is separated into groups or strata
 - must have an adequate number of events in each stratum (at least 5 to 7)
 - can be used to adjust for variables with strong impact on survival
 - can help solve proportional hazards violations
- Strata have different baseline hazards
- Coefficients / Hazard Ratios are calculated within stratum then combined.
- Vittinghoff 6.3.2

```
Session 8:
   Survival
analysis part 3
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```

formula

Call:

##

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How to stratify Example - in R, strata() can be added to any model

```
mycox <- coxph(Surv(time, os) ~ trt + strata(stage),</pre>
                data = pbc.os)
summary(mycox)
```

0.558

1.283

p = 0.6

p = 0.6

p = 0.6

```
## coxph(formula = Surv(time, os) ~ trt + strata(stage), data = ;
```

n= 312, number of events= 125

coef exp(coef) se(coef) z Pr(>|z|)## trt -0.1063 0.8992 0.1814 -0.586

exp(coef) exp(-coef) lower .95 upper .95 ## 1.112 0.6302 ## trt 0.8992 ##

Concordance= 0.494 (se = 0.025) ## Likelihood ratio test= 0.34 on 1 df.

Wald test = 0.34 on 1 df. ## Score (logrank) test = 0.34 on 1 df,

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What are competing risks?

- Example from Vittinghoff 6.5: The MrOS study (Orwoll et al. 2005) followed men over 65 to examine predictors of bone fracture and low BMD (subclinical bone loss)
- At end of study participants had:
 - developed fracture (outcome of interest),
 - remained alive without fracture (incomplete follow-up), or
 - died prior to fracture (incomplete follow-up)

Orwoll, E. et al. (2005). Design and baseline characteristics of the osteoporotic fractures in men (MrOS) study–a large observational study of the determinants of fracture in older men. Contemporary Clinical Trials, 26(5), 569–585.

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Why not treat died prior to fracture and alive without fracture as censored?

- Recall the independent censoring assumption (Vittinghoff 6.6.4):
 - censored people are similar to those who remain at risk in terms of developing the event of interest;
 - censoring is independent of the event of interest.
 - For patients who died this assumption is highly suspect

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Reasons for right censored data

- Cut-off date of analysis (administrative censoring):
 - Censoring usually independent
- Loss to follow-up
 - Independence may be problematic if sicker individuals discontinue participant in study (lack of energy, too ill, return to home country)
 - or if healthier individuals discontinue participation (don't feel the need to continue, start new life in other country)
- Competing risks:
 - Often informative.
 - In competing risks analysis, independence between competing risks is not required

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Very brief summary of competing risk methods

- 1-to-1 mapping between hazard and cumulative incidence function is lost in competing risks
- Standard Kaplan-Meier estimator is biased for competing risks data
 - Aalen-Johansen estimator is better choice
- Gary's test is analogous to log-rank test
- cause-specific standard Cox PH model might be useful for prognostic (causal) testing, but not estimating a population Hazard Ratio

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Resources for competing risk methods

- Z. Zhang, Survival analysis in the presence of competing risks, Ann Transl Med. 2017 Feb; 5(3): 47. PMID: 28251126
- cmprsk package
- riskRegression package

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What is propensity score analysis?

 an alternative to multivariate regression to control for hypothesized confounders in observational studies:

outcome ~ exposure + counfounder1 + confounder2

- a stratification approach that is more practical than stratifying on multiple hypothesized confounders
- an approach to summarizing many covariates into a single score
- a convenient approach to controlling for many hypothesized confounders

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Propensity score approach to correction for confounders

• *Step 1*: fit the propensity score model (no outcome) that predicts propensity for exposure based on confounders:

exposure ~ counfounder1 + confounder2

- Step 2: use propensity predictions to match or stratify participants with similar propensity (for example, stratifying on quintiles of propensity)
- *Step 3*: check adequacy of matching or stratification, ie by comparing attributes of matched participants
- Step 4: test hypothesis among matched participants:

outcome ~ exposure

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Propensity score references

- P.C. Austin (2011), An Introduction to Propensity Score Methods for Reducing the Effects of Confounding in Observational Studies. Multivariate Behavioral Research, 46:3, 399-424, DOI: 10.1080/00273171.2011.568786
- R. d'Agostino (1998), Tutorial in Biostatistics: propensity score methods for bias reduction in the comparison of a treatment to a non-randomized control group. Stat. Med. 17, 2265-2281. http:

 $//www.stat.ubc.ca/\sim john/papers/DAgostinoSIM1998.pdf$

 You don't need any special package to do basic propensity score matching (e.g. stratifying by quintiles), but the MatchIt package provides multiple matching approaches, diagnostics, good documentation