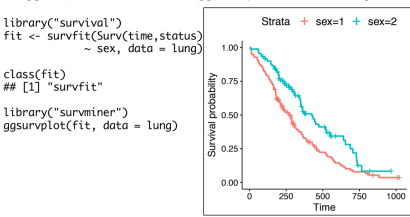
Creating Survival Plots

Informative and Elegant with *survminer*

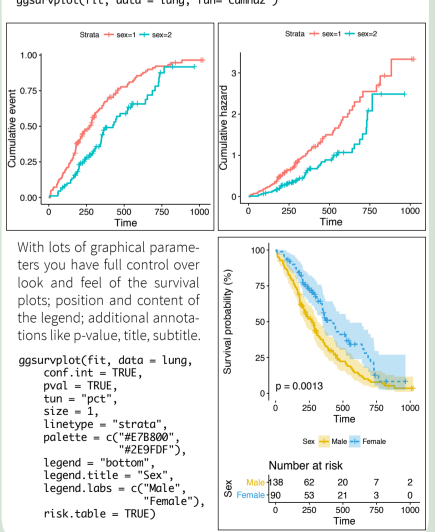
Survival Curves

The **ggsurvplot()** function creates **ggplot2** plots from **survfit** objects.



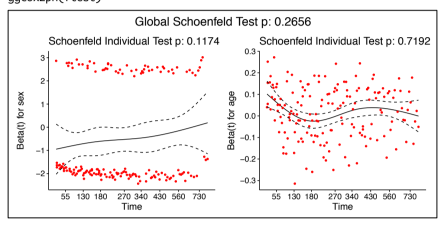
Use the **fun** argument to set the transformation of the survival curve. E.g. **"event"** for cumulative events, **"cumhaz"** for the cumulative hazard function or **"pct"** for survival probability in percentage.

```
ggsurvplot(fit, data = lung, fun="event")
ggsurvplot(fit, data = lung, fun="cumhaz")
```



Diagnostics of Cox Model

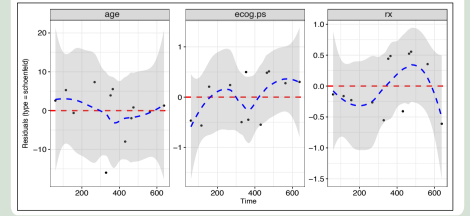
The function **cox.zph()** from **survival** package may be used to test the proportional hazards assumption for a Cox regression model fit. The graphical verification of this assumption may be performed with the function **ggcoxzph()** from the **survminer** package. For each covariate it produces plots with scaled Schoenfeld residuals against the time.



The function **ggcoxdiagnostics()** plots different types of residuals as a function of time, linear predictor or observation id. The type of residual is selected with **type** argument. Possible values are "martingale", "deviance", "score", "schoenfeld", "dfbeta", "dfbetas", and "scaledsch".

The **ox.scale** argument defines what shall be plotted on the OX axis. Possible values are "linear.predictions", "observation.id", "time".

Logical arguments **hline** and **sline** may be used to add horizontal line or smooth line to the plot.



Summary of Cox Model

The function **ggforest()** from the **survminer** package creates a forest plot for a Cox regression model fit. Hazard ratio estimates along with confidence intervals and p-values are plotter for each variable.

```
library("survival")
library("survminer")
lung$age <- ifelse(lung$age > 70, ">70","<= 70")</pre>
fit <- coxph( Surv(time, status) ~ sex + ph.ecog + age, data = lung)</pre>
## coxph(formula = Surv(time, status) ~ sex+ph.ecog+age, data=lung)
##
##
              coef exp(coef) se(coef)
## sex
                        0.567
                                 0.168 -3.37 0.00075
            -0.567
## ph.ecog 0.470
                        1.600
                                 0.113 4.16 3.1e-05
## aae>70
            0.307
                       1.359
                                 0.187
                                         1.64 0.10175
                                               Forest plot for coxph model
## Likelihood ratio test=31.6 on
## n= 227, number of events= 164
                                                     0.57 (p.value 0.00075***)
ggforest(fit)
                                                              1.6 (p.value 0.000031*
                                         ph.ecoa
                                                                 1.4 (p.value 0.1)
                                         age>70
```

The function **ggcoxadjustedcurves()** from the **survminer** package plots Adjusted Survival Curves for Cox Proportional Hazards Model. Adjusted Survival Curves show how a selected factor influences survival estimated from a Cox model.

Note that these curves differ from Kaplan Meier estimates since they present expected ssurvival based on given Cox model.

library("survival")
library("survminer")

lung\$sex <- ifelse(lung\$sex == 1, "Male", "Female")

fit <- coxph(Surv(time, status) ~ sex + ph.ecog + age, data = lung)

ggcoxadjustedcurves(fit, data=lung, variable=lung\$sex)

variable=lung\$sex)

variable Male Female

1.00
0.75
0.05
0.00
0 250 500 750 1000

Note that it is not necessary to include the grouping factor in the Cox model. Survival curves are estimated from Cox model for each group

c(35,55,65,85))

defined by the factor independently.

lung\$age3 <- cut(lung\$age,</pre>

ggcoxadjustedcurves(fit, data=lung, variable=lung\$age3)

variable (35,55] (55,65] (65,85]

Hazard ratio