# LDA Interpretation

# January 2018

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## Introduction

Summary of models and especially their interpretation (graphically as well as content based) used in Survival Analysis. This document emerged throughout the exam preparation for a lecture on Survival Data Analysis at LMU in winter 2018. Most examples are based on that lecture taught by Prof. Kuechenhoff and Andreas Bender.

## Censoring

#### 1. Right:

- 1. Type 1: study ends before event occured. E.g.: fixed time study of 1 year
- 2. Type 2: ...
- 3. Type 3: person withdraws from study because of other event. E.g.: interest on cancer death, person is getting shot
- 2. **Left**: we know when event occurs but we do not know when it started. E.g.: person dies at week 4 on cancer but we don't know the time of the disease outbreak. Our observed survival time of 4 weeks is thus equal (best case) or smaller then the observed.
- 3. **Left Truncation**: Biased because only people that survived made it to the study. E.g.: deductible in insurances, people with losses < deductibles are not getting observed.

## Kaplan Meier

#### **Model Equation**

Estimate the **Survival rate** non-parametrically without any covariables:

$$\hat{S}(t) = \prod_{t_k \le t} (1 - d_k/n_k), \forall t \ge t_1$$

where  $d_k$  = number of events at time point  $t_k$  (neither dead nor censored) and  $n_k =$  amount of people under risk right before time  $t_k$ .

Reveals a step function with jumps at each  $t_k$  where events took place.

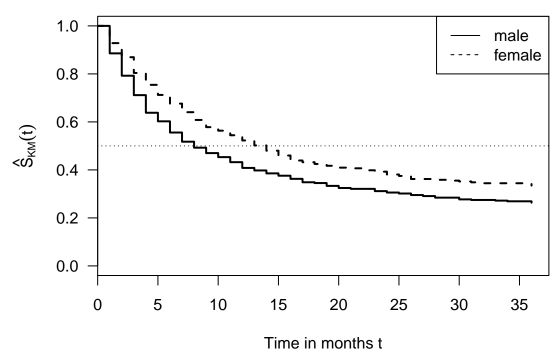
#### Data

This is some random SOEP data and we estimate Survival functions for both genders:

##		dauer	status	beginn.monat	female	${\tt male}$	alter	bild
##	1	11	0	114	0	1	47	1
##	2	30	1	83	0	1	38	2
##	3	1	1	83	0	1	44	2
##	4	36	0	85	0	1	28	2
##	5	1	1	111	0	1	38	2
##	6	7	0	104	1	0	30	1

#### Model

# Duration of unemployment by gender (Kaplan–Meier estimator)



#### Test

```
## Call:
## survdiff(formula = Surv(dauer, status) ~ female, data = soep)
##
##
               N Observed Expected (O-E)^2/E (O-E)^2/V
## female=0 1206
                      726
                                651
                                         8.62
                                                   22.1
## female=1 794
                      396
                               471
                                        11.92
                                                   22.1
   Chisq= 22.1 on 1 degrees of freedom, p= 2.6e-06
```

#### Nelson Aalen

#### Accelerated Failure Time Transformation models

Assumptions: 1. covariates have a multiplicative effect on the **Survival time**. E.g.: Survival time for smokers is an accelerated version of the survival time for non-smokers.

2. the survival time follows an assumed distribution that we get applying a density transformation

We model the survival time directly with the log-trafo:

$$log(T) = Y = \beta_0 + X^T \beta + \sigma \epsilon$$

$$T = exp(Y) = exp(\beta_0) * exp(X^T \beta) * exp(\sigma \epsilon)$$
 with  $\epsilon \sim$  Distribution e.g.: SEV, Normal, logistic, .....

Thus, the effect of the estimated coefficient  $\hat{\beta}_j$  on Survival time T is  $exp(\beta_j)$ 

Steps for the estimation: 1. calculate density for T 2. classic Maximum Likelihood Estimation

#### Examples for all AFT's

The exponential and the weibull AFT can be compared with a Cox PH model as they also have proportional (time independent) hazard ratios. This means that the following re-parametrization holds:

$$\beta_{PH} = -\frac{\beta_{\text{AFT: WB or Exp}}}{\sigma}$$

where sigma is our scale parameter from the AFT model equation.

Therefore we compare with this baseline Cox model:

```
## Call:
## coxph(formula = Surv(futime, fustat) ~ ecog.ps + rx, data = ovarian)
##
##
     n= 26, number of events= 12
##
              coef exp(coef) se(coef)
                                            z Pr(>|z|)
                      1.4474
## ecog.ps 0.3698
                               0.5869 0.630
                                                 0.529
                               0.5878 -0.984
## rx
           -0.5782
                      0.5609
                                                 0.325
##
##
           exp(coef) exp(-coef) lower .95 upper .95
## ecog.ps
              1.4474
                         0.6909
                                    0.4582
                                               4.573
## rx
              0.5609
                         1.7829
                                    0.1772
                                               1.775
##
## Concordance= 0.622 (se = 0.088)
## Rsquare= 0.054
                    (max possible= 0.932)
## Likelihood ratio test= 1.45
                                on 2 df,
                                            p=0.4833
## Wald test
                        = 1.43
                                on 2 df,
                                            p=0.4897
## Score (logrank) test = 1.46 on 2 df,
                                            p=0.4808
```

#### Exponential

```
##
## Call:
## survreg(formula = Surv(futime, fustat) ~ ecog.ps + rx, data = ovarian,
##
       dist = "exponential")
##
                Value Std. Error
                                       z
                           1.322 5.267 1.39e-07
## (Intercept)
                6.962
               -0.433
                           0.587 -0.738 4.61e-01
## ecog.ps
                           0.587 0.991 3.22e-01
## rx
                0.582
##
## Scale fixed at 1
##
## Exponential distribution
## Loglik(model) = -97.2
                          Loglik(intercept only) = -98
## Chisq= 1.67 on 2 degrees of freedom, p= 0.43
```

```
## Number of Newton-Raphson Iterations: 4
## n= 26
```

#### **AFT** interpretation:

- Geometric mean of survival time: 1055
- 1 unit change in ecog.ps shortens survival time by  $\exp(-0.433) = 0.65$
- 1 unit change in rx increases survival time by  $\exp(0.582) = 1.79$
- though, both effects are non significant

```
## (Intercept) ecog.ps rx
## 1055.5715021 0.6484732 1.7887244
```

#### PH interpretation (invert the coefficients)

- 1 unit change in ecog.ps increases the hazard h(t) by  $1/\exp(0.433) = 1.54$
- 1 unit change in rx decreases h(t) by 0.56

```
## (Intercept) ecog.ps rx
## 0.0009473541 1.5420838523 0.5590576235
```

#### Weibull

```
##
## Call:
## survreg(formula = Surv(futime, fustat) ~ ecog.ps + rx, data = ovarian,
##
      dist = "weibull")
##
                Value Std. Error
## (Intercept) 6.897
                           1.178 5.857 4.72e-09
                           0.527 -0.731 4.65e-01
              -0.385
## ecog.ps
                0.529
                           0.529 0.999 3.18e-01
## rx
## Log(scale) -0.123
                           0.252 -0.489 6.25e-01
## Scale= 0.884
##
## Weibull distribution
                          Loglik(intercept only)= -98
## Loglik(model) = -97.1
## Chisq= 1.74 on 2 degrees of freedom, p= 0.42
## Number of Newton-Raphson Iterations: 5
## n= 26
```

#### **AFT** interpretation:

- Geometric mean of survival time: 988
- 1 unit change in ecog.ps shortens survival time by  $\exp(-0.385) = 0.68$
- 1 unit change in rx increases survival time by  $\exp(0.529) = 1.70$
- though, both effects are non significant

- If scale parameter {R, echo = FALSE} survregWB\$scale was close to 1 we would yield an exponential model. Our shape parameter is 1 / scale
- coefficients: {R, echo = FALSE} exp(coef(survregWB))

#### PH interpretation (multiply by -1 and the shape parameter before exp())

- 1 unit change in ecog.ps increases the hazard h(t) by 1.55
- 1 unit change in rx decreases h(t) by 0.55

```
## (Intercept) ecog.ps rx
## 0.0004085855 1.5459383069 0.5498547398
```

#### Log Normal

Only AFT Interpretation!

- 1 unit incrase in ecog.ps shortens survival time by  $\exp(-.229) = 0.79$
- 1 unit incrase in rx increases survival time by  $\exp(0.813) = 2.25$
- Can we interpret the scale parameter?
- MORE TO ADD! DISCUSS

```
##
## Call:
## survreg(formula = Surv(futime, fustat) ~ ecog.ps + rx, data = ovarian,
##
       dist = "lognormal")
##
                Value Std. Error
                                      z
## (Intercept) 5.878
                           1.094 5.373 7.72e-08
## ecog.ps
               -0.229
                           0.537 -0.427 6.70e-01
## rx
                0.813
                           0.537 1.514 1.30e-01
                0.167
                           0.228 0.731 4.65e-01
## Log(scale)
##
## Scale= 1.18
## Log Normal distribution
## Loglik(model) = -95.9
                          Loglik(intercept only) = -97.1
## Chisq= 2.35 on 2 degrees of freedom, p= 0.31
## Number of Newton-Raphson Iterations: 3
## n = 26
```

#### Log logistic

```
##
## Call:
## survreg(formula = Surv(futime, fustat) ~ ecog.ps + rx, data = ovarian,
       dist = "loglogistic")
##
##
                Value Std. Error
                                       Z
                           1.134 5.435 5.49e-08
## (Intercept)
               6.161
## ecog.ps
               -0.336
                           0.537 -0.626 5.32e-01
                0.705
                           0.539 1.308 1.91e-01
## rx
                           0.248 -1.466 1.43e-01
## Log(scale) -0.363
##
## Scale= 0.695
##
## Log logistic distribution
```

```
## Loglik(model)= -96.3 Loglik(intercept only)= -97.4
## Chisq= 2.07 on 2 degrees of freedom, p= 0.36
## Number of Newton-Raphson Iterations: 4
## n= 26
```

#### Weibull Model

#### Model equation

model Survival time directly

$$T = exp(Y) = exp(\beta_0) * exp(X^T \beta) * exp(\sigma \epsilon)$$
 with  $\epsilon \sim$  Distribution e.g.: SEV, Normal, logistic, .....

Then use density-transformation to get distribution for T (Survival time).

Steps: 1. calculate density for T 2. classic Maximum Likelihood Estimation

#### Data

where delta depicts the event indicator (delta = 1: non-censored, delta = 0: censored)

```
type time delta
##
## 1
         1
              1
## 2
              3
                      1
         1
## 3
              3
         1
                      1
## 4
              4
         1
                      1
## 5
         1
             10
                      1
## 6
         1
             13
                      1
```

#### Model

```
##
## Call:
## survreg(formula = Surv(time, delta) ~ as.factor(type), data = tongue,
       dist = "weibull")
##
                     Value Std. Error
##
                                          7.
## (Intercept)
                     4.972
                                0.227 21.93 1.46e-106
## as.factor(type)2 -0.669
                                0.351 -1.91 5.66e-02
## Log(scale)
                     0.216
                                0.116 1.87 6.21e-02
##
## Scale= 1.24
##
## Weibull distribution
## Loglik(model) = -298.9
                           Loglik(intercept only) = -300.7
## Chisq= 3.58 on 1 degrees of freedom, p= 0.059
## Number of Newton-Raphson Iterations: 5
```

- Survival time decreases by factor  $\exp(-0.669) = 0.512$  for patient with type 2 w.r.t. type 1 given constant other features
- Not significant with p = 0.056
- Failure time accelerates by factor 1 / 0.512 = 1.953125 DISCUSS

- Scale Parameter = 1.24 -> if it was 1.0 we would yield an exponential distributed model. This would yield  $\lambda t = \lambda$ , constant and independent of time.
- Weibull model: not constant  $\lambda(t) = \lambda * \alpha(\lambda * t)^{\alpha-1}$  but hazard ratio still independent of time -> PH assumption

## Cox Regression model

Estimates coefficents  $\beta$  that have multiplicative effect on time-dependent hazard  $\lambda_0(t)$ . The baseline hazard is estimated non-parametrically via Breslow estimate. Thus, we yield step-functions for visualization, estimation,

super sweet R-bloggers post on Cox models

#### Model equation

$$\lambda_i(t) = \lambda_0(t) exp(x_i'\beta)$$

To get the estimator for the cumulative Hazard and the Survival rate:

- 1. estimate  $\beta$ s via Cox
- 1. estimate non-parametrically baseline hazards  $\lambda_0(t)$  e.g. via Breslow
- 1. ^^^calculate for each t  $\lambda(t) = \lambda_0(t) exp(x_i'\beta)$
- 1. cumulate the  $\lambda(t)$  to the cumulative Hazards  $\Lambda_t = \sum_{i=1}^t \lambda_i$
- 1. calculate estimated Survival  $S(t) = exp(-\Lambda_t)$

#### Data

where delta depicts the event indicator (delta = 1: non-censored, delta = 0: censored)

```
type time delta
## 1
         1
               1
                      1
## 2
               3
         1
                      1
## 3
               3
                      1
         1
## 4
         1
               4
                      1
## 5
         1
              10
                      1
## 6
         1
              13
                      1
```

#### Model

We are searching for the effect of the binary treatment type.

- Person with type 2 has a multiplicative factor  $\exp(0.4664) = 1.594245$  higher hazard rate than a person with type 1 (ceteris paribus in case of other covariates)
- this effect is not significant as the H0 can not be rejected at  $\alpha = 0.05$ , REMIND but this does not imply testing of the PH assumption
- log rank score test: tests for significant differencies in the survival curves for the two subpopulations seperated by the variable of interest (here: treatment). This means that the probability of an event occurring at any time point is the same for each subpopulation. H0: they do not differ -> p > 0.05: H0 cannot be rejected -> no significant effect of treatment. WHAT HAPPENS WITH MORE COVARIATES? E.G.: one significant, the other not

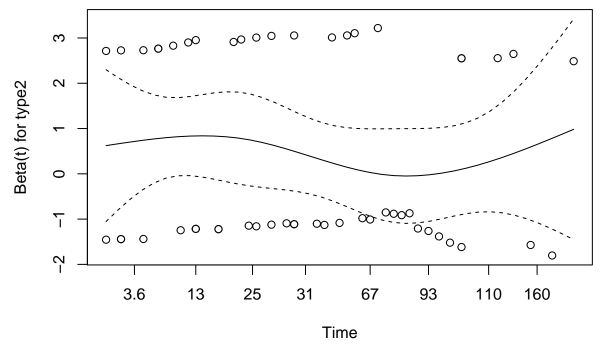
Summary of the Cox-PH model:

```
## Call:
## coxph(formula = Surv(time, delta) ~ type, data = tongue)
##
##
     n= 80, number of events= 53
##
##
           coef exp(coef) se(coef)
                                        z Pr(>|z|)
  type2 0.4664
##
                   1.5942
                             0.2804 1.663
##
                           0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
  Signif. codes:
##
##
         exp(coef) exp(-coef) lower .95 upper .95
  type2
             1.594
                       0.6273
                                  0.9201
                                              2.762
##
##
## Concordance= 0.564
                       (se = 0.036)
## Rsquare= 0.033
                     (max possible= 0.993 )
## Likelihood ratio test= 2.67
                                 on 1 df,
                                            p=0.102
## Wald test
                         = 2.77
                                 on 1 df,
                                            p=0.09632
## Score (logrank) test = 2.81
                                 on 1 df,
                                            p=0.09343
```

#### Test the Cox PH assumption for the covariates

#### Graphically

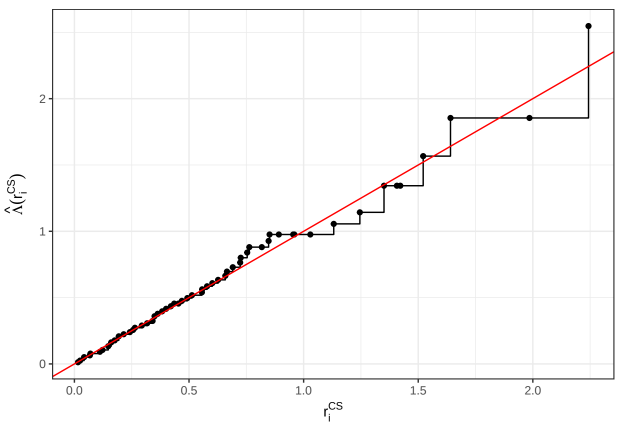
The scaled Schoenfeld residuals are used for that test and plotted against the time. Do this for each covariate to check the PH assumption for each covariate. If they **randomly and unstructured** center around zero: PH assumption holds! If not, not. The plot estimates a smooth function of the residuals over time for better visualization. Holds here:



Test PH Also based on Schoenfeld residuals, not exam-relevant. If p >> 0.05 there is no violation of the PH.

#### Test overall fit

Plot Cox-Snell residuals vs. Cumulated Hazard. If they share the diagnonal, everything is fine and we have a good overall model fit.

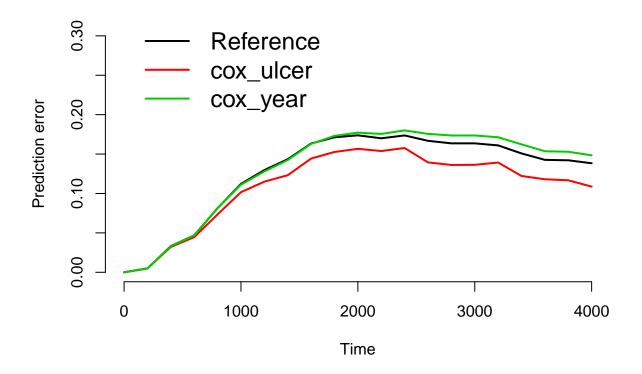


## Model fit Analysis

## Prediction Error Curves (PEC)

The predicted survival time for each time point is compared with the true survival time within the **Brier Score**. Some magic is added such as *inverse probability of censoring weights (IPCW)* to account for right censoring. Then scores for each time point are computed using Cross-Validation and the Brier Scores over time are plotted for all desired models. The lower the score, the better. This method is **model agnostic**.

For Melanoma compare predictive performance of Cox model with only variable ulcer as predictor with the reference Kaplan-Meier estimates and a Cox-PH model that uses year as a linear predictor. We see, that our cox-model outperforms the simple Kaplan-Meier estimator (which does not use any variables) and both outperform the stupid Cox model with time as linear predictor.



#### Residuals

- Schoenfeld
- Martingale
- Deviance
- Cox-Snell

#### Schoenfeld Residuals

Use case: test PH assumption for each covariate

Idea: compute Schoenfeld residuals for Variable k and m observations. Those residuals should be independent of the survival time. This is the test that cox.zph() performs.

#### Test

##		rho	chisq	р
##	fin	0.0267	0.0838	0.77227
##	age	-0.2264	7.5618	0.00596
##	prio	-0.0657	0.5330	0.46533
##	mar	0.1327	2.1143	0.14593
##	employed.lag1	-0.0427	0.2066	0.64942
##	GLOBAL	NA	9.4135	0.09366

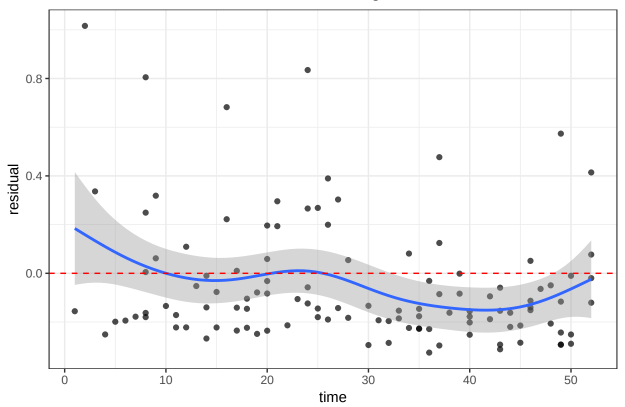
Small p-value for variable age indicates problem with the PH assumption here. High value for employed.lag1 indicates nice fulfillment of PH assumption.

Can we observe this graphically?

#### Graphically

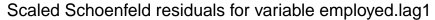
Plot the Schoenfeld residuals for variable age:

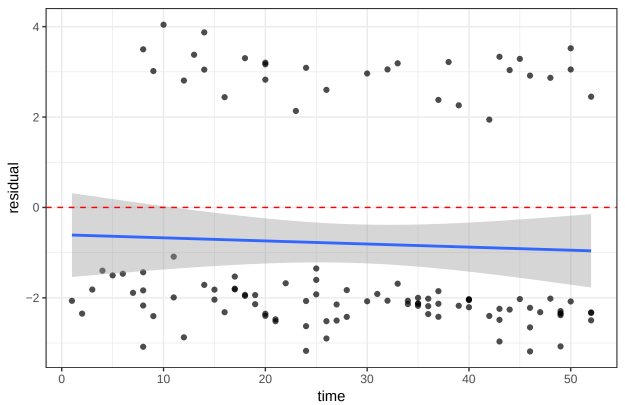
## Scaled Schoenfeld residuals for variable age



PH assumption violated because there is non linear structure in the data.

What can we do? Exclude variable or model time varying effect // non-linearly e.g. using splines Check variable employed lag1 that had huge p-value in zph test (good sign for PH):





We see what we expected: there seems to be no PH violation. Sweet!

#### Cox Snell residuals

Use case: Check overall goodness of fit

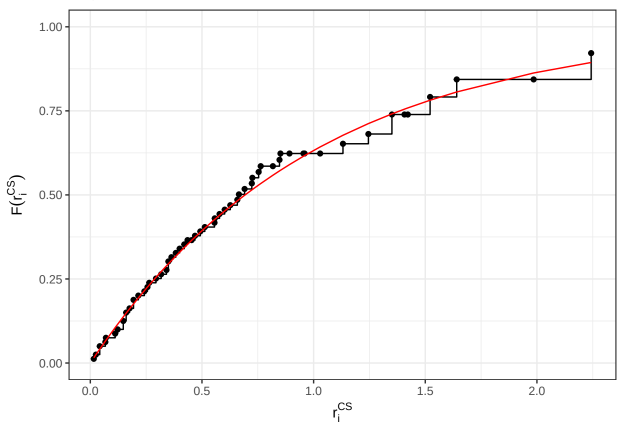
#### Model

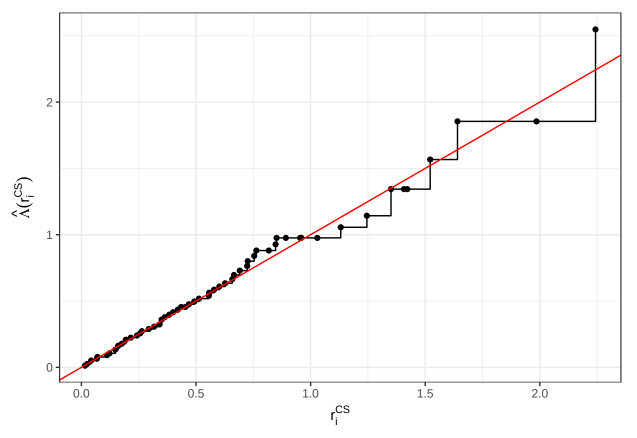
#### Example

Check the overall goodness of fit for a simple cox model:

```
## coxph(formula = Surv(time, delta) ~ type, data = tongue)
##
    n= 80, number of events= 53
##
           coef exp(coef) se(coef)
##
                                      z Pr(>|z|)
## type2 0.4664
                   1.5942
                            0.2804 1.663
                                          0.0963 .
##
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
         exp(coef) exp(-coef) lower .95 upper .95
##
             1.594
                                0.9201
## type2
                       0.6273
## Concordance= 0.564 (se = 0.036)
```

```
## Rsquare= 0.033 (max possible= 0.993 )  
## Likelihood ratio test= 2.67 on 1 df, p=0.102  
## Wald test = 2.77 on 1 df, p=0.09632  
## Score (logrank) test = 2.81 on 1 df, p=0.09343
```





Two options: 1. Plot cs-residuals against estimated distribution Function values. Their distribution should then follow a standard exponential distribution if the model is fit correctly. 2. Plot against estimated cumulative hazard function. This should result in a straight line if the model fits the data.

## Semi-parametric additive Cox model

## Time discrete Survival models

## Piecewise exponential models (PEM)

#### Model equation:

$$\lambda_i(t|x_i) = \lambda_j exp(x^T \beta), \forall t \in ]a_{j-1}, a_j]$$

with constant baseline hazards in each of the J intervals.

#### Data

##		id	tstart	tend	interval	${\tt offset}$	${\tt ped\_status}$	${\tt treatment}$	pair
##	1	1	0	1	(0,1]	0	1	placebo	1
##	2	2	0	1	(0,1]	0	0	6-MP	1
##	3	2	1	2	(1,2]	0	0	6-MP	1
##	4	2	2	3	(2,3]	0	0	6-MP	1

```
We fit an intercept-only model for many intervals resulting in many baseline intercepts:
##
## Call:
  glm(formula = ped_status ~ interval - 1, family = poisson(link = log),
##
       data = leuk.ped, offset = offset)
##
##
   Deviance Residuals:
##
       Min
                 10
                      Median
                                    30
                                            Max
                     -0.3162
##
   -0.7559
            -0.3780
                               -0.2294
                                          2.3082
##
  Coefficients:
##
                    Estimate Std. Error z value Pr(>|z|)
##
## interval(0,1]
                      -3.0445
                                  0.7071
                                          -4.306 1.67e-05 ***
                     -2.9957
                                  0.7071
                                          -4.237 2.27e-05 ***
## interval(1,2]
## interval(2,3]
                      -3.6376
                                  1.0000
                                          -3.638 0.000275 ***
## interval(3,4]
                      -2.9178
                                  0.7071
                                          -4.126 3.69e-05 ***
## interval(4,5]
                      -2.8622
                                  0.7071
                                          -4.048 5.17e-05 ***
## interval(5,6]
                     -2.3979
                                  0.5774
                                          -4.153 3.28e-05 ***
                      -3.3673
## interval(6,7]
                                  1.0000
                                          -3.367 0.000759 ***
## interval(7,8]
                      -1.9459
                                  0.5000
                                          -3.892 9.95e-05 ***
## interval(8,9]
                     -19.3026
                               1924.2001
                                           -0.010 0.991996
## interval(9,10]
                      -3.1355
                                  1.0000
                                          -3.135 0.001716 **
## interval(10,11]
                      -2.3514
                                  0.7071
                                          -3.325 0.000883 ***
                      -2.1972
## interval(11,12]
                                  0.7071
                                           -3.107 0.001888 **
                                          -2.773 0.005561 **
## interval(12,13]
                     -2.7726
                                  1.0000
## interval(13,15]
                     -3.4012
                                  1.0000
                                          -3.401 0.000671 ***
## interval(15,16]
                      -2.6391
                                  1.0000
                                          -2.639 0.008314 **
## interval(16,17]
                      -2.5649
                                  1.0000
                                           -2.565 0.010319 *
## interval(17,19]
                    -19.9957
                               2842.2319
                                          -0.007 0.994387
## interval(19,20]
                    -19.3026
                               2980.9580
                                           -0.006 0.994833
## interval(20,22]
                      -2.1972
                                  0.7071
                                           -3.107 0.001888 **
## interval(22,23]
                     -1.2528
                                  0.7071
                                           -1.772 0.076449
## interval(23,25]
                    -19.9957
                               4215.7112
                                          -0.005 0.996216
## interval(25,32]
                    -21.2485
                               4713.3084
                                           -0.005 0.996403
## interval(32,34]
                    -19.9957
                               6665.6247
                                           -0.003 0.997606
## interval(34,35]
                    -19.3026
                               9426.6168
                                          -0.002 0.998366
##
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
##
   (Dispersion parameter for poisson family taken to be 1)
##
       Null deviance: 1017.84 on 475 degrees of freedom
## Residual deviance: 148.11
                               on 451 degrees of freedom
##
  AIC: 256.11
##
## Number of Fisher Scoring iterations: 17
```

• we fit way too many parameters

## 5

## 6 2

3

4

4

5

(3,4]

(4,5]

0

6-MP

6-MP

1

1

- intervals which did not face events have super high standard errors and strange coefficients
- -> Two reasons for fitting PAM's with smooth baseline hazards

## Piecewise additive exponential models (PAM)

New compared to PEM: smooth modeling of the piecewise constant baseline hazards e.g. via splines. Cool because:

- PEM constrained by use of intervals as high J leads to parameter explosion
- Smoother curves due to penalization of splines on the overlaps of the intervals
- Problem PEM: no data in interval  $]a_{l-1}, a_l] \rightarrow \lambda_l = 0$ , wiggely hazard rate curves

#### Model equation:

$$\lambda_i(t|x_i) = exp(f_0(t_i) + x^T \beta)$$

with spline for time dependent baseline hazard:

$$f_0(t_j) = log(\lambda_0(t_j)) = \sum_{k=1}^{K} \gamma_k B_k(t_j)$$

and for time varying covariates:

$$\lambda_i(t|x_i) = exp(f_0(t_j) + \sum_{j=1}^{p} f_k(x_i, k))$$

## Piecewise additive exponential mixed models (PAMM)

#### Model equation:

$$\lambda_i(t|x_i) = exp(f_0(t_i) + x^T \beta)$$

with spline for time dependent baseline hazard:

$$f_0(t_j) = log(\lambda_0(t_j)) = \sum_{k=1}^{K} \gamma_k B_k(t_j)$$

and for time varying covariates:

$$\lambda_i(t|x_i) = exp(f_0(t_j) + \sum_{i=1}^{p} f_k(x_i, k))$$

#### Data

looks like that:

##		CombinedI	) tstart	tend	interval	${\tt offset}$	ped_status	${\tt CombinedicuID}$	Year	Age
##	1	110:	1 4	5	(4,5]	0	0	1114	2007	71
##	2	110:	1 5	6	(5,6]	0	0	1114	2007	71
##	3	110:	1 6	7	(6,7]	0	0	1114	2007	71
##	4	110:	1 7	8	(7,8]	0	0	1114	2007	71
##	5	110:	1 8	9	(8,9]	0	0	1114	2007	71
##	6	110:	1 9	10	(9,10]	0	0	1114	2007	71
##		BMI A	AdmCatID	DiagI	D2 Apache	eIIScore	e DaysInICU			
##	1	38.97392	${\tt Medical}$	Seps	is	13	6.743056			

```
## 2 38.97392 Medical Sepsis 13 6.743056

## 3 38.97392 Medical Sepsis 13 6.743056

## 4 38.97392 Medical Sepsis 13 6.743056

## 5 38.97392 Medical Sepsis 13 6.743056

## 6 38.97392 Medical Sepsis 13 6.743056
```

Fit a PAMM with a smooth spline term for time (tend) and the other continuous variables using this formula:

```
pamm_icu <- bam(ped_status ~ s(tend) + Year + AdmCatID + DiagID2 + s(Age) + s(BMI) +
        s(ApacheIIScore) + s(CombinedicuID, bs="re"), offset=offset, data = ped,
        family=poisson(), discrete = TRUE)</pre>
```

We include the variable CombinedicuID as a random effect aka as a **frailty term**. Therefore wie use bs = "re". We control for the random effects of the ICU units without having to model a dummy for each of the ICU's. The frailty model just estimates a Gaussian over the different ICU's for which we only have to estimate the variance: 1 parameter vs. 400.

We model the PAM as a Poisson model with log link on the death-indicator ped\_status

This is the model summary:

```
##
## Family: poisson
## Link function: log
## Formula:
## ped_status ~ s(tend) + Year + AdmCatID + DiagID2 + s(Age) + s(BMI) +
       s(ApacheIIScore) + s(CombinedicuID, bs = "re")
##
##
## Parametric coefficients:
##
                              Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                                          0.11388 -40.383 < 2e-16 ***
                              -4.59863
## Year2008
                               0.02718
                                          0.07425
                                                    0.366 0.714314
## Year2009
                              -0.08622
                                          0.07466
                                                  -1.155 0.248156
## Year2011
                              -0.02329
                                          0.06966
                                                  -0.334 0.738144
## AdmCatIDSurgical Elective
                             -0.47450
                                          0.09297
                                                   -5.104 3.33e-07 ***
## AdmCatIDSurgical Emergency -0.25668
                                          0.07228
                                                  -3.551 0.000384 ***
## DiagID2Cardio-Vascular
                               0.12439
                                          0.08721
                                                    1.426 0.153774
                                          0.12855
## DiagID2Other
                               0.10391
                                                    0.808 0.418914
## DiagID2Metabolic
                              -0.92768
                                          0.25552
                                                  -3.631 0.000283 ***
## DiagID2Neurologic
                               0.01267
                                          0.09508
                                                    0.133 0.893972
## DiagID2Orthopedic/Trauma
                              -0.26816
                                          0.11560
                                                  -2.320 0.020354
## DiagID2Renal
                              -0.02734
                                          0.21580
                                                  -0.127 0.899183
## DiagID2Respiratory
                              -0.13289
                                          0.08618
                                                  -1.542 0.123091
## DiagID2Sepsis
                               0.05627
                                          0.09895
                                                    0.569 0.569587
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Approximate significance of smooth terms:
##
                        edf
                             Ref.df Chi.sq p-value
## s(tend)
                      1.000
                              1.001 248.94
                                           < 2e-16 ***
                      1.002
                              1.003 122.98 < 2e-16 ***
## s(Age)
## s(BMI)
                              3.879 40.61 3.55e-08 ***
                      3.061
## s(ApacheIIScore)
                      1.890
                              2.422 163.17 < 2e-16 ***
## s(CombinedicuID) 101.279 363.000 152.16 3.35e-08 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

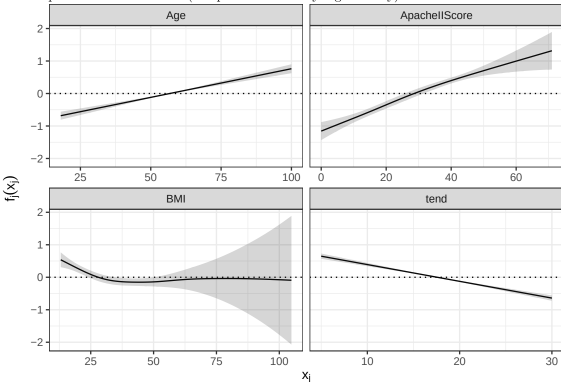
```
##
## R-sq.(adj) = -0.00897 Deviance explained = -15%
## fREML = 2.0196e+05 Scale est. = 1 n = 208536
```

#### What can we say?

- smooth terms for continuos variables:
  - if the edf (estimated degress of freedom) = 1, our spline smoother estimated the variable as a linear effect on the hazard rate. This is the case for Age and time
  - BMI, ApacheIIScore and CombinedicuID (only frailty effect) seem to have a non-linear effect on the hazard rate

#### - HOW TO INTERPRET SPLINE FOR COMBINEDICUID FRAILTY @ ANDREAS

- those effects can also be seen graphically which shows the effect of the variable's values on the linear predictor aka the log(hazard-rate). This is the exact value that enters our linear predictor, e.g. 75 year old person enters 0.3
- time (tend) has a falling slope aka a decreasing effect on the log(hazard) -> has hazard decreases also
- ApacheIIScore has almost linear effect: (log-) hazard increases with increasing Apache Scores though this increase is getting lower with higher values of the score
- increasing linear age effect, the older, the higher the (log-)hazard
- typical shape of the BMI effect, very low BMIs have increased hazard, that decreases toward "normal" BMIs, high uncertainty with respect to effect of very high BMIs as number of patients with respective BMIs decreases (few persons with very high obesity)



- non-smooth terms for categorical variables:
  - exponentiate the coefficients exp(beta) and interpret their mulitplicative effect on the hazard

rate w.r.t the reference category

- example 1: hazard rate for a person treated in 2009 is  $\exp(-0.08622441) = 0.9173883$  times as high as the hazard rate for similar person treated in 2007 (reference category)
- example 2: hazard rate for a person with Metabolic cancer is  $\exp(-0.92767602) = 0.3954717$  times as high as the hazard rate for similar person with Gastrointestinal cancer (reference category)
- For more, interpret this table:

##		beta	HR
##	Year2008	0.02718222	1.0275550
##	Year2009	-0.08622441	0.9173883
##	Year2011	-0.02328905	0.9769801
##	AdmCatIDSurgical Elective	-0.47449956	0.6221964
##	${\tt AdmCatIDSurgical\ Emergency}$	-0.25667793	0.7736173
##	DiagID2Cardio-Vascular	0.12438947	1.1324568
##	DiagID2Other	0.10391129	1.1095020
##	DiagID2Metabolic	-0.92767602	0.3954717
##	DiagID2Neurologic	0.01267184	1.0127525
##	DiagID2Orthopedic/Trauma	-0.26815998	0.7647854
##	DiagID2Renal	-0.02733998	0.9730304
##	DiagID2Respiratory	-0.13289109	0.8755604
##	DiagID2Sepsis	0.05627062	1.0578839

### Frailty models

#### Aalen model

model equation

$$\lambda(t) = \lambda_0(t) + x'(t)\beta(t) = \sum_{k=1}^{p} x_k(t)\beta_k(t)$$

with additive effects of time-varying covariates on baseline hazard rate

### Cox-Aalen model

#### model equation

$$\lambda(t) = \lambda_0(t) + X(t)\beta(t) \cdot exp(Z(t)'\gamma)$$

with additive effects of time-varying covariates on baseline hazard rate which are also multiplicatively affected via Cox part of the model.  $\gamma$  are time-constant coefficients, PH-assumption, and  $\beta$  are time varying additive coefficients by the Aalen-part.

#### Data

looks like that

```
##
     major_complications age charlson_score sex transfusion metastasesYN
## 1
                            58
                                                   f
                        no
                                              2
                                                              yes
## 2
                       yes
                            52
                                              2
                                                               no
                                                                              1
## 3
                                              2
                                                   f
                                                                              1
                            74
                        no
                                                              yes
                                              2
## 4
                       yes
                            57
                                                   m
                                                              yes
                                                                              1
## 5
                            30
                                              2
                                                   f
                                                                              1
                        no
                                                              yes
## 6
                                              2
                                                   f
                        no
                                                              yes
                                                                              1
##
     major_resection days status id metastases
## 1
                        579
                                     1
## 2
                                  0
                                     2
                   no 1192
                                               yes
## 3
                        308
                                     3
                                               yes
                   no
## 4
                         33
                                     4
                  yes
                                               yes
## 5
                        397
                                  1
                                     5
                  yes
                                               yes
## 6
                  yes 1219
                                  0
                                     6
                                               yes
```

#### What can we say from the graphic?

#### • Age:

- the cumulative Hazard of a person aged A+1 at time point t=1500 is 0.01 higher than that of a person aged A
- the effect of metastases on the cumulative hazard rate starts to increase t = 1000 after the surgery and is approx. constant before

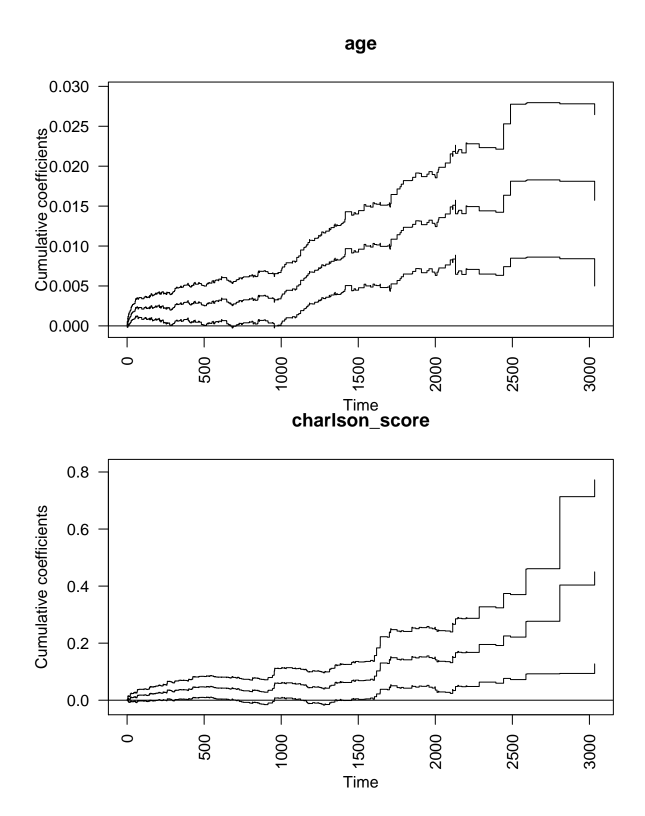
#### • Complications:

- the cumulative Hazard of a person with major complications at time point t = 1500 is 0.2 higher than that of a person without complications
- the effect of complications on the cumulative hazard rate decreases over time

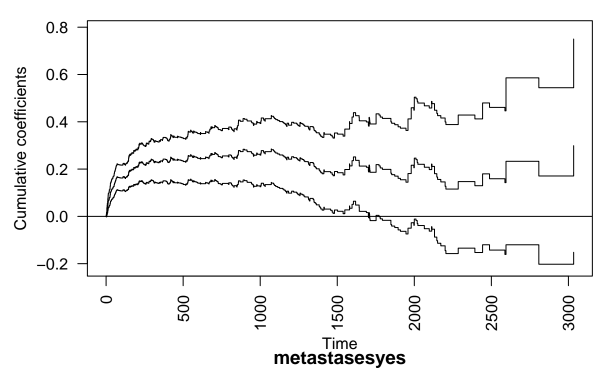
#### • Metastases:

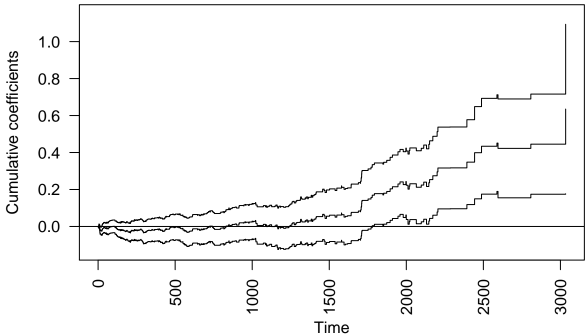
- the cumulative Hazard of a person with metastases at time point t=2500 is 0.4 higher than that of a person without metastases
- the effect of metastases on the cumulative hazard rate starts to matter only after t = 1500 and then increases more or less linearly
- before t = 1500 the effect is non significant as the 0 is part of the confidence intervals

Effects for the continuous variables estimated as additive via the Aalen-part of the model using the formula Surv(days, status) ~ age + charlson\_score + major\_complications + metastases + prop(sex) + prop(transfusion) + prop(major\_resection), data = liver, residuals = 1, basesim = 1)



## major\_complicationsyes





What can we say from the model summary?

## Cox-Aalen Model

##

## Test for Aalen terms

```
## Test for nonparametric terms
##
## Test for non-significant effects
                           Supremum-test of significance p-value H_0: B(t)=0
## (Intercept)
                                                     4.00
                                                                         0.002
                                                     4.18
                                                                         0.002
## age
## charlson score
                                                     4.04
                                                                         0.000
## major_complicationsyes
                                                     6.07
                                                                         0.000
## metastasesyes
                                                     3.85
                                                                         0.000
##
## Test for time invariant effects
##
                                 Kolmogorov-Smirnov test
## (Intercept)
                                                  0.43700
## age
                                                  0.00522
## charlson_score
                                                  0.16400
## major_complicationsyes
                                                  0.21200
## metastasesyes
                                                  0.28100
##
                           p-value H_0:constant effect
## (Intercept)
                                                  0.190
## age
                                                  0.410
## charlson_score
                                                  0.060
## major_complicationsyes
                                                  0.138
## metastasesyes
                                                  0.020
##
## Proportional Cox terms :
                                      SE Robust SE D2log(L)^-1
##
                             Coef.
                                                                   z P-val
## prop(sex)f
                             0.224 0.111
                                             0.107
                                                          0.109 2.08 0.0373
## prop(transfusion)yes
                                                          0.112 2.07 0.0386
                             0.233 0.111
                                              0.113
## prop(major_resection)yes 0.254 0.113
                                                          0.113 2.31 0.0207
                                             0.110
##
                             lower2.5% upper97.5%
## prop(sex)f
                               0.00644
                                            0.442
## prop(transfusion)yes
                               0.01540
                                            0.451
## prop(major_resection)yes
                               0.03250
                                            0.475
## Test of Proportionality
                                  hat U(t) | p-value H 0
## prop(sex)f
                                         9.53
                                                      0.178
## prop(transfusion)yes
                                         6.51
                                                      0.568
## prop(major_resection)yes
                                         8.99
                                                      0.198
```

- Aalen part:
  - Supremum-test: for all 4 variables the H0: no effect can be rejected
  - Kolmogorov Smirnov for time variant effects: H0: constant effect can only clearly be rejected for metastases DISCUSS THIS
- Cox part:
  - sexf: the additive, time-varying effects  $\beta(t) = (\beta_{age}(t), \beta_{charlson}(t), \beta_{complications}(t), \beta_{metastases}(t))^T$  from the Aalen model is getting multiplied by factor exp(0.224) = 1.251071 for a female compared with a similar man
  - same for transfusion ( $\exp(0.233) = 1.262381$ ) and major resection ( $\exp(0.254) = 1.289172$ )
  - DISCUSS

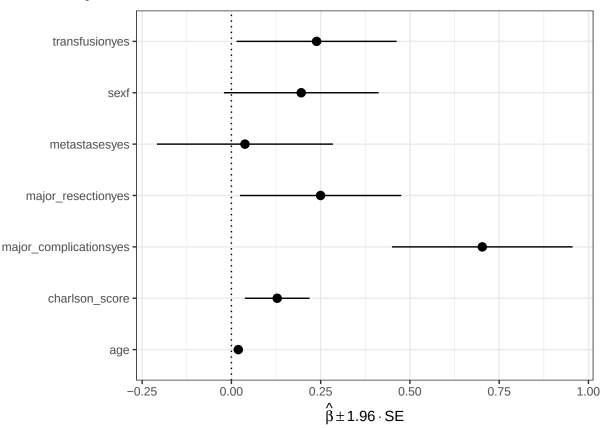
#### Cox-Aalen vs. PAM

Compare this with the PAM fitted on the data using the below formula. We explicitly model time varying effects of the 4 variables (metastases, marjo complications, age, charlson) as in the Aalen model via ti().

```
bam(
  formula = ped_status ~ ti(tend,k=10) +
    # use ti() for non-identifiability issue
    metastases + ti(tend, by = as.ordered(metastases),k=10, mc = c(1,0)) +
    major_complications + ti(tend,by = as.ordered(major_complications),k=10, mc = c(1,0)) +
    age + ti(tend, by = age,k=10, mc = c(1,0)) +
    charlson_score + ti(tend, by = charlson_score,k=10, mc = c(1,0)) +
    sex + transfusion + major_resection,
    data = ped_liver,
    offset = offset,
    family = poisson())
```

The figure below shows the effect of the time constant variables which allow some interpretation:

- NOTE: Constant contributions to time-varying can be interpreted as effects at t=0. Check the model equation and DISCUSS
- sex: Compared to males, females have a 1.22 times increased risk of experiencing an event (c.p.)
- transfusion: Compared to patients without transfusion, patients with transfusion have a 1.27 times increased risk of experiencing an event (c.p.)
- major resection: A major resection increases the risk of event by a factor of 1.28, compared to patients without a major resection
- DISCUSS If above interpretation holds, this would fit nicely the effect of the time-constant factors in the Cox-part of above Cox-Aalen model

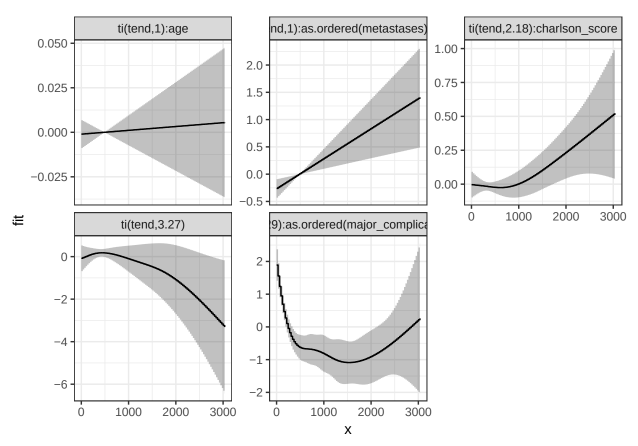


Model summary:

##
## Family: poisson
## Link function: log

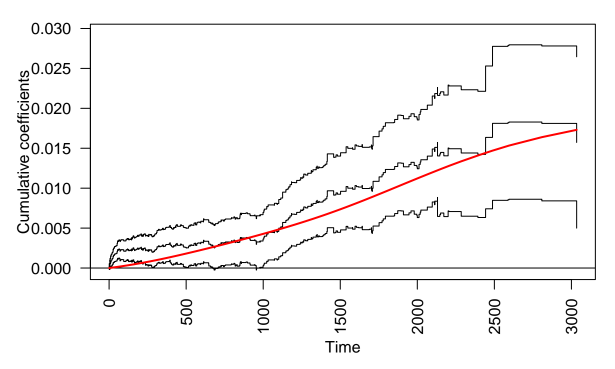
```
##
## Formula:
  ped_status ~ ti(tend, k = 10) + metastases + ti(tend, by = as.ordered(metastases),
      k = 10, mc = c(1, 0)) + major_complications + ti(tend, by = as.ordered(major_complications),
##
       k = 10, mc = c(1, 0)) + age + ti(tend, by = age, k = 10,
      mc = c(1, 0)) + charlson_score + ti(tend, by = charlson_score,
##
      k = 10, mc = c(1, 0)) + sex + transfusion + major_resection
##
##
## Parametric coefficients:
##
                           Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                          -9.756319
                                     0.384061 -25.403 < 2e-16 ***
                                     0.123233
                                                0.308 0.758122
## metastasesyes
                           0.037949
## major_complicationsyes
                          0.702678
                                     0.126452
                                                 5.557 2.75e-08 ***
                           0.019308
                                     0.005269
                                                 3.664 0.000248 ***
                                                 2.833 0.004604 **
## charlson_score
                           0.128265
                                      0.045268
## sexf
                           0.195558
                                      0.108301
                                                 1.806 0.070967 .
                                                 2.128 0.033311 *
## transfusionyes
                           0.238512
                                      0.112066
## major_resectionyes
                           0.249730
                                      0.112940
                                                 2.211 0.027024 *
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Approximate significance of smooth terms:
##
                                                 edf Ref.df Chi.sq p-value
## ti(tend)
                                               3.266 3.960 9.103
                                                                    0.05775
## ti(tend):as.ordered(metastases)yes
                                               1.003 1.005 9.513 0.00208
## ti(tend):as.ordered(major_complications)yes 5.289 6.165 70.698 5.55e-13
## ti(tend):age
                                               1.000 1.001 0.068 0.79468
## ti(tend):charlson_score
                                               2.183 2.682 7.672 0.05013
##
## ti(tend)
## ti(tend):as.ordered(metastases)yes
## ti(tend):as.ordered(major_complications)yes ***
## ti(tend):age
## ti(tend):charlson_score
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## R-sq.(adj) = 0.000679
                           Deviance explained = -10.1%
## fREML = 2.7942e+05 Scale est. = 1
                                              n = 147896
```

This is the effect estimated for the smooth terms. The total effect of x at time point t is  $\beta_x * x + f_x(t)$  where  $\beta_x * x$  are the constant effects from the previous graphic and  $f_x(t)$  models the effect of the smooth time varying term. Recap the PAM model equation  $\lambda_i(t|x_i) = exp(f_0(t_j) + x^T\beta)$  and DISCUSS. They look like that:

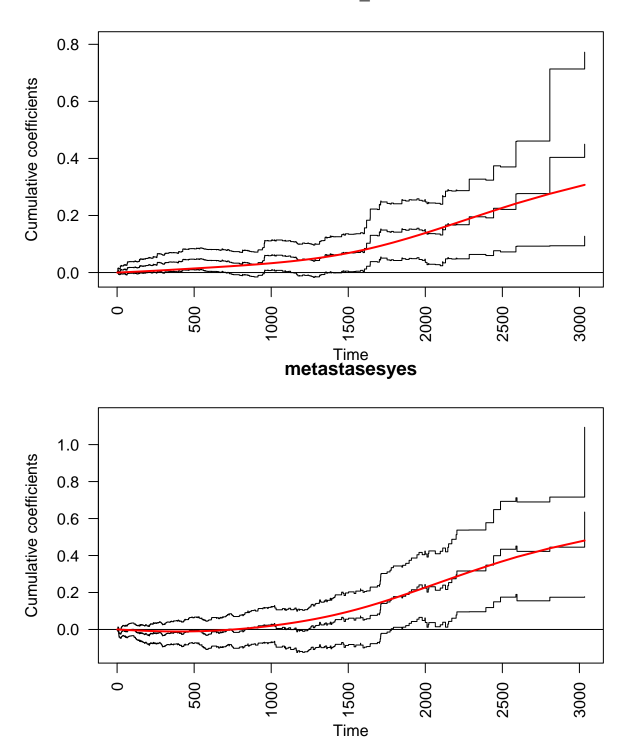


Visual comparison of the time-varying effects from Cox-Aalen model on the cumulated Hazard over time (black) vs. the smooth multiplivative effects of the PAM model (red).

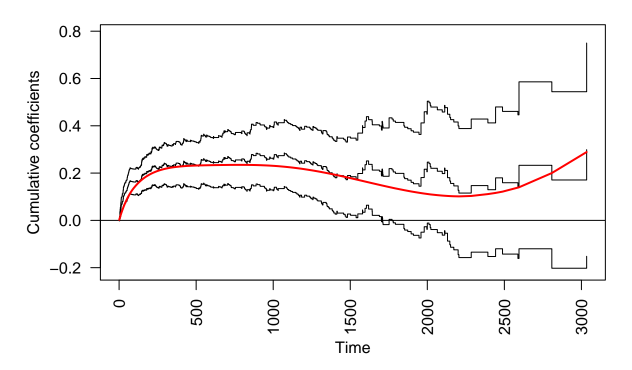




# charlson\_score



## major\_complicationsyes



## Competing Risk models

## Random Stuff

Attest significance only based on  $\beta$  and  $se(\beta)$ 

- 1. compute z-score:  $z = \beta/se(\beta)$
- 2. thresholds for alpha=0.05:
- 3. one sided:  $z_{thresh} = 1.64$
- 4. two sided:  $z_{thresh} = 1.96$
- 5. Reject H0 (coefficient is not significant) if  $z>z_{thresh}$

check their explanation