Introduction to the rstpm2 package

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Abstract

This vignette outlines the methods and provides some examples for generalised survival models as implemented in the R rstpm2 package.

Keywords: survival, splines.

1. Background and theory

Generalised survival models provide a flexible and general approach to modelling survival or time-to-event data. The survival function S(t|x) to time t for covariates x is defined in terms of a inverse link function G and a linear prediction $\eta(t,x)$, such that

$$S(t|x;\theta) = G(\eta(t,x;\theta))$$

where η is a function of both time t and covariates x, with regression parameters θ . We can calculate the hazard from this function, where

$$h(t|x; \theta) = \frac{d}{dt} \left(-\log(S(t|x; \theta)) \right)$$
$$= \frac{-G'(\eta(t, x; \theta))}{G(\eta(t, x; \theta))} \frac{\partial \eta(t, x; \theta)}{\partial t}$$

We model using a linear predictor $\eta(t, x; \theta) = X(t, x)\theta$ for a design matrix X(t, x). The linear predictor can be constructed in a flexible manner, with the main constraint being that the time effects be smooth and twice differentiable. We calculate the derivative for the linear predictor using finite differences, such that

$$\frac{\partial \eta(t, x; \theta)}{\partial t} = \frac{\partial X(t, x)\theta}{\partial t} = \frac{X(t + \epsilon, x) - X(t - \epsilon, x)}{2\epsilon} \theta = X_D(t, x)\theta$$

for a derivative design matrix $X_D(t,x)$. This formulation allows for considerable flexibility in the construction of the linear predictor, with possible interactions between time and covariates.

The default smoother for time using natural splines for $\log(\text{time})$, which is the flexible parametric survival model developed by Royston and Parmar (2003) and implemented by the Stata command stpm2 1

The models are estimated using maximum likelihood estimation (MLE) for fully parametric models, penalised MLE for penalised smoothers, maximum marginal likelihood estimation

¹As a technical aside, the Stata implementation uses natural splines using a truncated power basis with orthogonalisation, while the ns() function in R uses a matrix projection of B-splines. Note that we have implemented an extended nsx() function for natural splines that includes cure splines, centering, and a compatibility argument to use Stata stpm2's unusual specification of quantiles.

Link description	Inverse link function $G(\eta(t, x; \theta))$	Interpretation	link.type
log-log	$\exp(-\exp(\eta(t,x;\theta)))$	Proportional hazards	"PH"
logit	$\operatorname{expit}(-\eta(t,x; heta))$	Proportional odds	"P0"
probit	$\Phi(-\eta(t,x; heta))$	Probit	"probit"
\log	$\exp(-\eta(t,x;\theta))$	Additive hazards	"AH"
Aranda-Ordaz	$\exp(-\log(\psi * \exp(\eta(t, x; \theta)) + 1)/\psi)$	Aranda-Ordaz	"AO"

Table 1: Implemented link functions

(MMLE) for parametric models with clustered data, or penalised MMLE for penalised models with clustered data. The likelihoods include left truncation, right censoring and interval censoring. For clustered data, we include Gamma frailties and normal random effects. Details on these models are available from https://doi.org/10.1177/0962280216664760 and https://doi.org/10.1002/sim.7451.

2. Syntax

The main functions for fitting the models are stpm2 for parametric models, possibly with clustered data, and pstpm2 for penalised models, possibly with clustered data. A subset of the syntax for stpm2 is:

```
stpm2(formula, data, smooth.formula = NULL,
    df = 3, tvc = NULL,
    link.type=c("PH","PO","probit","AH","AO"), theta.AO=0,
    bhazard = NULL,
    robust = FALSE, cluster = NULL, frailty = !is.null(cluster) & !robust,
    RandDist=c("Gamma","LogN"),
    ...)
```

The formula has a Surv object on the left-hand-side and a linear predictor on the right-hand-side that does not include time (for pstpm2, it also does not include penalised functions). The time effects can be specified in several ways: the most general is using smooth.formula, where the right-hand-side of the formula specifies functions for time that are smooth with respect to time. This specification can include interactions between time and covariates. As an example, smooth.formula=~nsx(log(time),df=3)+x:nsx(log(time),df=2) specifies a baseline natural spline smoother of the log of the variable time used in the Surv object with three degrees of freedom, with an interaction between a covariate x and a natural spline smoother of log(time) with two degrees of freedom. Other specifications of time effects have equivalent formulations: for example, df=3 is equivalent to smooth.formula=~nsx(log(time),df=3) for the variable time. Similarly, tvc=list(x=2) is equivalent to smooth.formula=~x:nsx(log(time),df=2). Moreover, for a log-linear interaction between a covariate and time, use smooth.formula=~x:log(time)

Type of link is specified with the link.type argument; this defaults to a log-log link for proportional hazards (see Table 1). For the Aranda-Ordaz link, the fixed value of the scale term ψ is specified using the theta.AO argument. For relative survival, a vector for the baseline hazard can be specified using the bhazard argument. A vector for the clusters can be specified with the cluster argument. The calculation of robust standard errors can be specified with the robust=TRUE argument; if robust is false, then the model assumes a frailty or random effects model, with either a default Gamma frailty (RandDist="Gamma") or a normal random effect (RandDist="LogN", using notation from the frailtypack package).

A current limitation of the implementation is that the dataset data needs to be specified.

The default specification for the additive hazards (link.type=="AH") models follows that for the ahaz package on CRAN: for a model specified as

```
stpm2(Surv(time,event)~x, data=data, link.type="AH")
```

we assume natural splines for the baseline time effect and a constant hazard for a unit change in the covariate x; an equivalent specification is

```
stpm2(Surv(time,event)~1, data=data, link.type="AH", smooth.formula=~nsx(time,df=3)+x:time) where there is default smoother for time and an interaction between linear x and linear time.
```

The regression coefficient for x:time can be interpreted as the additive rate for a unit change in x.

The syntax for the fitting the penalised models with pstpm2 is very similar. A subset of the arguments are:

The penalised smoothers are specified using the s() function from the mgcv package within the smooth.formula argument; by default, not specifying smooth.formula will lead to

```
smooth.formula=~s(log(time))
```

Interactions with time (both penalised and unpenalised) and penalised covariate effects should be specified using smooth.formula. Note that the df argument is not included. By default, the smoothing parameter(s) are using the criterion argument; the smoothing parameters can also be fixed using the sp argument. The specifications for relative survival, link type, and clustered data follow that for the stpm2 function.

2.1. Post-estimation

One of the strengths of these models is varied post-estimation. Most of the estimators are described in Tables 2 and 3. These estimators are typically calculated from the predict function or from plot function calls. All of these calls require that the newdata argument is specified (in contrast to prediction in the survival package, which defaults to the average of each covariate).

For contrasts (e.g. survival differences, hazard ratios), the newdata argument is the "unexposed" group, while the exposed group is defined by either: (i) a unit change in a variable in newdata as defined by the var argument (e.g. var="x" for variable x); or (ii) an exposed function that takes a data-frame and returns the "exposed" group (e.g. exposed=function(data) transform(data, x=1)). The latter mechanism is quite general and allows for standardised survival, standardised hazards, and attributable fractions under possibly counterfactual exposures.

Standard errors for the post-estimators are calculated on a possibly transformed scale using the delta method. For the delta method, the partial derivatives of the post-estimators are calculated either directly or using finite differences.

Description	Formulation a	type
Conditional link	$\eta(t,x;\hat{ heta})$	"link"
Conditional survival	$S(t x;\hat{\theta}) = G(\eta(t,x;\hat{\theta}))$	"surv"
Conditional odds	$Odds(t x; \hat{\theta}) = S(t x; \hat{\theta}) / (1 - S(t x; \hat{\theta}))$	"odds"
Conditional failure	$1-S(t x;\hat{ heta})$	"fail"
Conditional cumulative hazard	$H(t x; \hat{\theta}) = -\log G(\eta(t, x; \hat{\theta}))$	"cumhaz"
Conditional density	$f(t x;\hat{\theta}) = G'(\eta(t,x;\hat{\theta})) \frac{\partial \eta(t,x;\theta)}{\partial t}$	"density"
Conditional hazard	$h(t x;\hat{\theta}) = \frac{G'(\eta(t,x;\hat{\theta}))}{G(\eta(t,x;\hat{\theta}))} \frac{\partial \eta(t,x;\hat{\theta})}{\partial t}$ $\log h(t x;\hat{\theta})$	"hazard"
Conditional log hazard	$\log h(t x;\hat{ heta})$	"loghazard"
Conditional survival differences	$S(t x^*; \hat{\theta}) - S(t x; \hat{\theta})$	"survdiff"
Conditional hazard differences	$h(t x^*;\hat{ heta}) - h(t x;\hat{ heta})$	"hazdiff"
Conditional hazard ratios	$h(t x^*;\hat{ heta})/h(t x;\hat{ heta})$	"hr"
Conditional odds ratios	$Odds(t x^*; \hat{\theta})/Odds(t x; \hat{\theta})$	"or"
Restricted mean survival time	$\int_0^t S(u x;\hat{\theta})du$	"rmst"
Standardised survival	$E_{X^*}S(t X^*;\hat{ heta})$	"meansurv"
Standardised survival differences	$E_{X_1^*}S(t X_1^*;\hat{\theta}) - E_{X_0^*}S(t X_0^*;\hat{\theta})$	"meansurvdiff"
Standardised hazard	$h_{X^*}(t X^*;\hat{\theta}) = \frac{E_{X^*}(\mathring{S}(t X^*;\hat{\theta})h(t X^*;\hat{\theta}))}{E_{X^*}(S(t X^*;\hat{\theta}))}$	"meanhaz"
Standardised hazard ratio	$h_{X_1^*}(t X_1^*;\hat{\theta})/h_{X_0^*}(t X_0^*\hat{\theta})$	"meanhazdiff"
Attributable fraction	$\frac{E_{X^*}S(t X^*;\hat{\theta}) - E_XS(t X;\hat{\theta})}{1 - E_XS(t X;\hat{\theta})}$	"af"

^aNotation: x^* is a covariate pattern for the "exposed" group; X^* is a set of possibly counterfactual covariates; $E_X(g(X))$ is the expectation or average of g(X) across the set X; X_0^* and X_1^* are sets of possibly counterfactual covariates for the "unexposed" and "exposed" sets, respectively.

Table 2: Types of conditional post-estimators

Description	Formulation a	type
Marginal survival	$S_M(t x;\hat{\theta}) = E_Z G(\eta(t,x,Z;\hat{\theta}))$	"margsurv"
Marginal hazard	$h_M(t x;\hat{\theta}) = E_Z(h(t,x,Z;\hat{\theta}))$	"marghaz"
Marginal survival differences	$S_M(t x^*;\hat{ heta}) - S_M(t x;\hat{ heta})$	"margsurvdiff"
Marginal hazard ratios	$h_M(t x^*;\hat{ heta})/h_M(t x;\hat{ heta})$	"marghr"
Standardised marginal survival	$E_Z E_{X^*} S(t X^*, Z; \hat{\theta})$	"meanmargsurv"
Standardised marginal survival differences	$E_Z E_{X^*} S(t X^*, Z; \hat{\theta}) - E_Z E_X S(t X, Z; \hat{\theta})$	"meansurvdiff"
Attributable fraction	$\frac{E_Z E_{X^*} S(t X^*,Z;\hat{\theta}) - E_Z E_X S(t X,Z;\hat{\theta})}{1 - E_Z E_X S(t X,Z;\hat{\theta})}$	"af"

[&]quot;Notation: Z is a random effect or frailty; x^* is a covariate pattern for the "exposed" group; X^* is a set of possibly counterfactual covariates; $E_X(g(X))$ is the expectation or average of g(X) across the set X; X_0^* and X_1^* are sets of possibly counterfactual covariates for the "unexposed" and "exposed" sets, respectively.

Table 3: Types of post-estimators for clustered data

3. Examples: Independent survival analysis

We begin with some simple proportional hazard models using the broancer dataset. We can fit the models using very similar syntax to coxph, except that we need to specify the degrees of freedom for the baseline smoother. Typical values for df are 3-6. For this model the model parameters include an intercept term, time-invariant log-hazard ratios, and parameters for the baseline smoother. The default for the baseline smoother is to use the nsx function, which is a limited extension to the splines::ns function, with log of the time effect.

Functionality	Uncorrelated param.	Uncorrelated penal.	Param. gamma frailty	Penal. gamma frailty	Param. normal random effects	Penal. normal random effects
Multiple links			\boxtimes			\boxtimes
Right censoring	\boxtimes	\boxtimes	\boxtimes			\boxtimes
Left truncation	\boxtimes	\boxtimes	\boxtimes^a	\boxtimes^a	\boxtimes^a	\boxtimes^a
Interval censoring	\boxtimes	\boxtimes				\boxtimes
Time-varying effects	\boxtimes	\boxtimes	\boxtimes	\boxtimes	\boxtimes	\boxtimes
Excess hazards	\boxtimes	\boxtimes	\boxtimes	\boxtimes	\boxtimes	\boxtimes
Conditional estimators b	\boxtimes	\boxtimes	\boxtimes	\boxtimes	\boxtimes	\boxtimes
Conditional standardisation c	\boxtimes	\boxtimes	\boxtimes	\boxtimes	\boxtimes	\boxtimes
Frailty/random effects variance			\boxtimes	\boxtimes	\boxtimes	\boxtimes
Marginal estimators d			\boxtimes	\boxtimes	\boxtimes	\boxtimes
Marginal standardisation e			\boxtimes	\boxtimes		
Random intercept			\boxtimes			\boxtimes
Random slope						\boxtimes
Multiple random effects						

 $[^]a{
m Gradients}$ not currently implemented.

Table 4: Functionality for the different generalised survival models

 $[^]b$ Estimators including survival, survival differences, hazards, hazard ratios, hazard differences, density, odds and odds ratios.

 $[^]c$ Standardised estimators include mean survival, mean survival differences, mean hazards and attributable fractions.

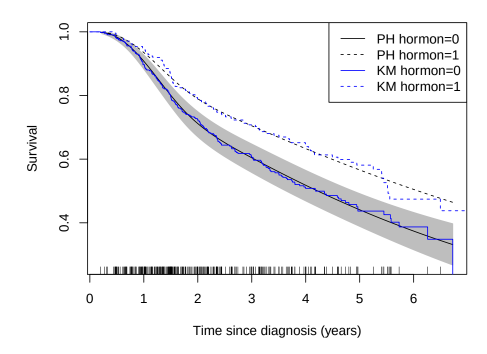
 $[^]d$ Marginal estimators include survival, survival differences, hazards, hazard ratios and hazard differences.

 $[^]e$ Marginal standardised estimators include mean survival, mean survival differences and attributable fractions.

```
> brcancer <- transform(brcancer, recyear=rectime / 365.24)</pre>
> fit <- stpm2(Surv(recyear,censrec==1)~hormon, data=brcancer, df=4)</pre>
> summary(fit)
Maximum likelihood estimation
Call:
stpm2(formula = Surv(recyear, censrec == 1) ~ hormon, data = brcancer,
   df = 4
Coefficients:
                          Estimate Std. Error z value
                                                         Pr(z)
(Intercept)
                          -6.79773 0.72642 -9.3578 < 2.2e-16 ***
hormon
                          nsx(log(recyear), df = 4)1 5.69995 0.71677 7.9523 1.830e-15 ***
nsx(log(recyear), df = 4)2 4.85614 0.48002 10.1166 < 2.2e-16 ***
nsx(log(recyear), df = 4)3 10.13328 1.41268 7.1731 7.331e-13 ***
nsx(log(recyear), df = 4)4 4.70626 0.33016 14.2545 < 2.2e-16 ***
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' '1
-2 log L: 1684.412
> ## utility
> eform.coxph <- function(object) exp(cbind(coef(object),confint(object)))</pre>
> fit.cox <- coxph(Surv(recyear,censrec==1)~hormon, data=brcancer)</pre>
> rbind(cox=eform(fit.cox),
        eform(fit)[2,,drop=FALSE])
                    2.5 %
                             97.5 %
hormon 0.6948843 0.5438441 0.8878726
hormon 0.6948513 0.5636725 0.8449349
```

We see that the hazard ratios are very similar to the coxph model. The model fit can also be used to estimate a variety of parameters. For example, we can easily estimate survival and compare with the Kaplan-Meier curves:

```
> plot(fit, newdata=data.frame(hormon=0), xlab="Time since diagnosis (years)")
> lines(fit, newdata=data.frame(hormon=1), lty=2)
> lines(survfit(Surv(recyear,censrec==1)~hormon, data=brcancer), col="blue", lty=1:2)
> legend("topright", c("PH hormon=0","PH hormon=1","KM hormon=0","KM hormon=1"),
+ lty=1:2, col=c("black","black","blue","blue"))
```

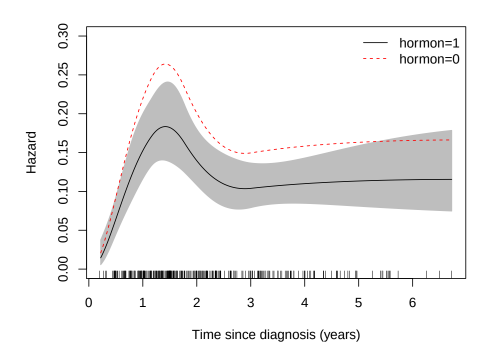


We can also calcu-

late the hazards.

```
> plot(fit,newdata=data.frame(hormon=1), type="hazard",
```

- + xlab="Time since diagnosis (years)", ylim=c(0,0.3))
- > lines(fit, newdata=data.frame(hormon=0), col=2, lty=2, type="hazard")
- > legend("topright", c("hormon=1","hormon=0"),lty=1:2,col=1:2,bty="n")

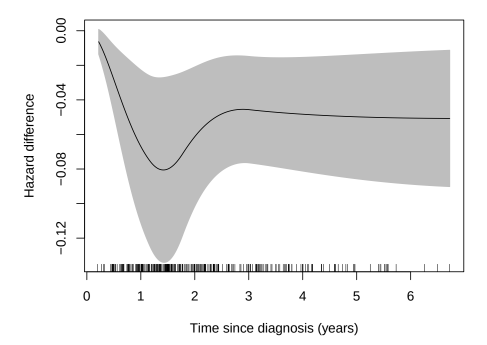


Usefully, we can also

estimate survival differences and hazard differences. We define the survival differences using a reference covariate pattern using the newdata argument, and then define an exposed function which takes the newdata and transforms for the 'exposed' covariate pattern. As an example:

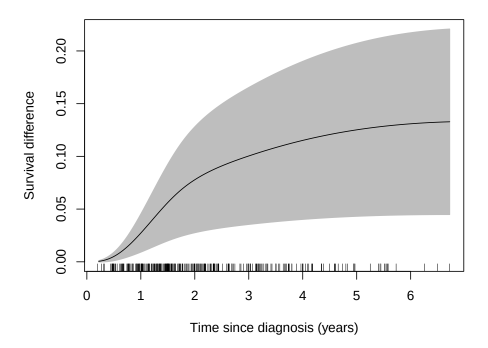
```
> plot(fit,newdata=data.frame(hormon=0), type="hdiff",
+ exposed=function(data) transform(data, hormon=1),
+ main="hormon=1 compared with hormon=0",
+ xlab="Time since diagnosis (years)")
```

hormon=1 compared with hormon=0



```
> plot(fit,newdata=data.frame(hormon=0), type="sdiff",
+ exposed=function(data) transform(data, hormon=1),
+ main="hormon=1 compared with hormon=0",
+ xlab="Time since diagnosis (years)")
```





4. Additive hazards

The additive hazards models takes the general form $H(t|x;\theta) = \eta(t,x;\theta)$. As a recent change, the default model specification for formula=Surv(t,e)~x without specifying smooth.formula is to use $H(t|x;\theta) = B(t)\theta_B + t(x^T\theta_x)$, where B(t) is a natural spline design matrix with parameters θ_B , and with θ_x as the parameters for x; the hazard is then $h(t|x;\theta) = B'(t)\theta_B + x^T\theta_x$.

The additive hazards have the attractive property that the effects are collapsible: adjusting for an uncorrelated covariate will not change the estimated conditional effect. These models have received some uptake within the causal inference field. This implementation is flexible, where the baseline (cumulative) hazard can be modelled using splines and we can model for both constant hazards and smooth effects over time. One possible issue with their interpretation is whether the rates will be approximately additive for different effects. One approach to conceptualise these models is to consider the effects as being competing risks and where we are adding competing risks together.

For our example using the breast cancer dataset with the randomised assignment to hormonal therapy, we find that hazard for those on hormonal therapy was -0.047 per year (95% confidence interval: -0.066, -0.024) compared with those not on hormonal therapy.

```
> brcancer <- transform(brcancer, recyear=rectime / 365.24)
> fit <- stpm2(Surv(recyear,censrec==1)~hormon, data=brcancer, link.type="AH")
> summary(fit)
```

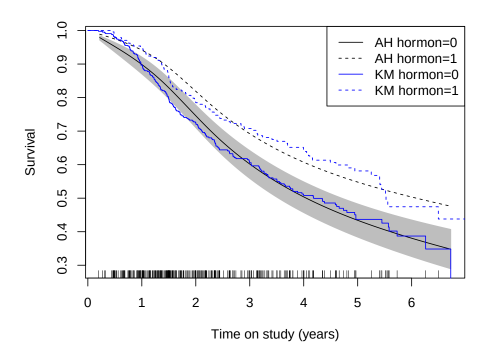
Maximum likelihood estimation

```
Call:
stpm2(formula = Surv(recyear, censrec == 1) ~ hormon, data = brcancer,
```

```
link.type = "AH")
```

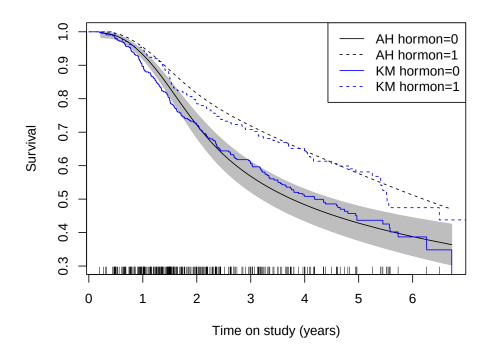
```
Coefficients:
```

```
Estimate Std. Error z value
                                                      Pr(z)
nsx(recyear, df = 3)1 0.597339
                                 0.047946 12.4585 < 2.2e-16 ***
nsx(recyear, df = 3)2 0.953091
                                 0.067957 14.0250 < 2.2e-16 ***
nsx(recyear, df = 3)3 0.984793
                                 0.082318 11.9632 < 2.2e-16 ***
hormon:recyear
                                  0.015486 -3.0122 0.002594 **
                     -0.046646
Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
-2 log L: 1733.751
> confint(fit)
                            2.5 %
                                       97.5 %
nsx(recyear, df = 3)1 0.53209270
                                  0.65983425
nsx(recyear, df = 3)2 0.92490181
                                  0.99227368
nsx(recyear, df = 3)3 0.92524890
                                   1.03931229
hormon:recyear
                      -0.06608299 -0.02365527
> plot(fit, newdata=data.frame(hormon=0), xlab="Time on study (years)")
> lines(fit, newdata=data.frame(hormon=1), lty=2)
> lines(survfit(Surv(recyear,censrec==1)~hormon, data=brcancer), col="blue", lty=1:2)
> legend("topright", c("AH hormon=0", "AH hormon=1", "KM hormon=0", "KM hormon=1"),
         lty=1:2, col=c("black","black","blue","blue"))
```



This can be modelled more flexibly using the **smooth.formula** argument. For example, we could model for **sqrt(recyear)** and include a natural spline smoother for the effect of **hormon**:

```
> fit <- stpm2(Surv(recyear,censrec==1)~1, data=brcancer, link.type="AH",
               smooth.formula=~ns(sqrt(recyear),df=3)+hormon:ns(recyear,df=3))
> summary(fit)
Maximum likelihood estimation
Call:
stpm2(formula = Surv(recyear, censrec == 1) ~ 1, data = brcancer,
    smooth.formula = ~ns(sqrt(recyear), df = 3) + hormon:ns(recyear,
        df = 3), link.type = "AH")
Coefficients:
                             Estimate Std. Error z value
                                                             Pr(z)
ns(sqrt(recyear), df = 3)1
                                        0.053216 12.0554 < 2.2e-16 ***
                             0.641539
ns(sqrt(recyear), df = 3)2
                             0.856835
                                        0.061732 13.8798 < 2.2e-16 ***
ns(sqrt(recyear), df = 3)3
                             1.005078
                                        0.086310 11.6450 < 2.2e-16 ***
hormon:ns(recyear, df = 3)1 -0.350244
                                        0.086176 -4.0643 4.818e-05 ***
hormon:ns(recyear, df = 3)2 -0.312674
                                        0.095885 -3.2609
                                                           0.00111 **
hormon:ns(recyear, df = 3)3 -0.244983
                                        0.128370 -1.9084
                                                           0.05634 .
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
-2 log L: 1705.101
> plot(fit, newdata=data.frame(hormon=0), xlab="Time on study (years)")
> suppressWarnings(lines(fit, newdata=data.frame(hormon=1), lty=2))
> lines(survfit(Surv(recyear,censrec==1)~hormon, data=brcancer), col="blue", lty=1:2)
> legend("topright", c("AH hormon=0","AH hormon=1","KM hormon=0","KM hormon=1"),
         lty=1:2, col=c("black","black","blue","blue"))
```



The square root trans-

form seems to considerably improve the fit at earlier times.

5. Mean survival

```
This has a useful interpretation for causal inference.
```

```
E_Z(S(t|Z,X=1)) - E_Z(S(t|Z,X=0)) fit <- stpm2(...) predict(fit,type="meansurv",newdata=data)
```

6. Cure models

For cure, we use the melanoma dataset used by Andersson and colleagues for cure models with Stata's stpm2 (see http://www.pauldickman.com/survival/).

Initially, we merge the patient data with the all cause mortality rates.

```
> popmort2 <- transform(rstpm2::popmort,exitage=age,exityear=year,age=NULL,year=NULL)
> colon2 <- within(rstpm2::colon, {</pre>
    status <- ifelse(surv_mm>120.5,1,status)
    tm <- pmin(surv_mm, 120.5)/12</pre>
    exit <- dx+tm*365.25
    sex <- as.numeric(sex)</pre>
    exitage <- pmin(floor(age+tm),99)</pre>
    exityear <- floor(yydx+tm)</pre>
    ##year8594 <- (year8594=="Diagnosed 85-94")
+ })
> colon2 <- merge(colon2,popmort2)</pre>
For comparisons, we fit the relative survival model without and with cure.
> fit0 <- stpm2(Surv(tm,status %in% 2:3)~I(year8594=="Diagnosed 85-94"),</pre>
+
                 data=colon2,
                 bhazard=colon2$rate, df=5)
>
> summary(fit <- stpm2(Surv(tm, status %in% 2:3)~I(year8594=="Diagnosed 85-94"),
                        data=colon2,
                        bhazard=colon2$rate,
                        df=5,cure=TRUE))
Maximum likelihood estimation
Call:
stpm2(formula = Surv(tm, status %in% 2:3) ~ I(year8594 == "Diagnosed 85-94"),
    data = colon2, df = 5, cure = TRUE, bhazard = colon2$rate)
Coefficients:
```

Estimate Std. Error z value

Pr(z)

```
(Intercept)
                                     -3.977319
                                                 0.054777 -72.6097 < 2.2e-16
I(year8594 == "Diagnosed 85-94")TRUE -0.155610
                                                 0.025088 -6.2026 5.554e-10
nsx(log(tm), df = 5, cure = TRUE)1
                                      3.323187
                                                 0.053164 62.5078 < 2.2e-16
nsx(log(tm), df = 5, cure = TRUE)2
                                      3.628625
                                                 0.053158 68.2608 < 2.2e-16
nsx(log(tm), df = 5, cure = TRUE)3
                                      1.634848
                                                 0.022464 72.7749 < 2.2e-16
nsx(log(tm), df = 5, cure = TRUE)4
                                      6.592010
                                                 0.111501
                                                           59.1206 < 2.2e-16
nsx(log(tm), df = 5, cure = TRUE)5
                                      3.371802
                                                 0.042787 78.8039 < 2.2e-16
(Intercept)
                                     ***
I(year8594 == "Diagnosed 85-94")TRUE ***
nsx(log(tm), df = 5, cure = TRUE)1
nsx(log(tm), df = 5, cure = TRUE)2
nsx(log(tm), df = 5, cure = TRUE)3
nsx(log(tm), df = 5, cure = TRUE)4
                                     ***
nsx(log(tm), df = 5, cure = TRUE)5
                                     ***
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
-2 log L: 42190.77
> predict(fit,head(colon2),se.fit=TRUE)
   Estimate
                lower
                          upper
1 0.8610828 0.8544630 0.8677538
2 0.7934654 0.7851997 0.8018182
3 0.6967405 0.6865297 0.7071032
4 0.8610828 0.8544630 0.8677538
5 0.8221244 0.8145060 0.8298140
6 0.8610828 0.8544630 0.8677538
```

The estimate for the year parameter from the model without cure is within three significant figures with that in Stata. For the predictions, the Stata model gives:

	+-			+
	 -	surv	surv_lci	surv_uci
1.		.86108264	.8542898	.8675839
2.	1	.79346526	.7850106	.8016309
3.		.69674037	.6863196	.7068927
4.		.86108264	.8542898	.8675839
5.		.82212425	.8143227	.8296332
	-			
6.	-	.86108264	.8542898	.8675839
	+-			+

We can estimate the proportion of failures prior to the last event time:

```
> newdata.eof <- data.frame(year8594 = unique(colon2$year8594),
+ tm=10)
> 1-predict(fit0, newdata.eof, type="surv", se.fit=TRUE)
```

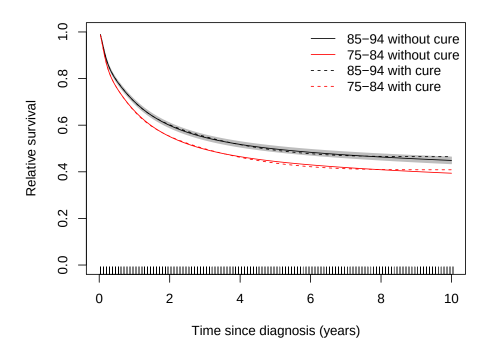
```
Estimate
                lower
                          upper
1 0.6060914 0.6205876 0.5910413
2 0.5512434 0.5655493 0.5364663
> 1-predict(fit, newdata.eof, type="surv", se.fit=TRUE)
  Estimate
                lower
                          upper
1 0.5913311 0.6052341 0.5769385
2 0.5350826 0.5482903 0.5214888
> predict(fit, newdata.eof, type="haz", se.fit=TRUE)
      Estimate
                      lower
                                   upper
1 1.254142.... 1.081724e-06 1.426561e-06
2 1.073411.... 9.235373e-07 1.223286e-06
We can plot the predicted survival estimates:
> tms=seq(0,10,length=301)[-1]
> plot(fit0,newdata=data.frame(year8594 = "Diagnosed 85-94", tm=tms), ylim=0:1,
       xlab="Time since diagnosis (years)", ylab="Relative survival")
> plot(fit0, newdata=data.frame(year8594 = "Diagnosed 75-84", tm=tms),
       add=TRUE,line.col="red",rug=FALSE)
> ## warnings: Predicted hazards less than zero for cure
> plot(fit,newdata=data.frame(year8594 = "Diagnosed 85-94",tm=tms),
       add=TRUE, ci=FALSE, lty=2, rug=FALSE)
```

> plot(fit,newdata=data.frame(year8594="Diagnosed 75-84",tm=tms),
+ add=TRUE,rug=FALSE,line.col="red",ci=FALSE,lty=2)
> legend("topright",c("85-94 without cure","75-84 without cure",

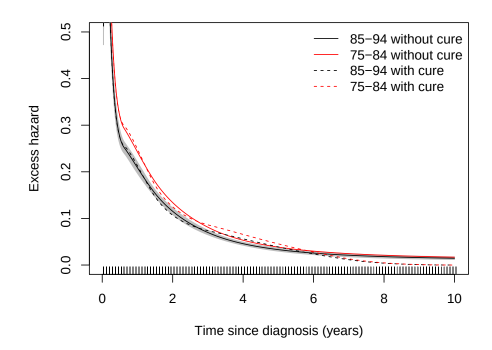
col=c(1,2,1,2), lty=c(1,1,2,2), bty="n")

"85-94 with cure", "75-84 with cure"),

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And the hazard curves:



7. Potential limitations and next steps

- The calculation of the matrix $X_D(t,x)$ using finite differences can be inaccurate for small event times. The functions stpm2 and pstpm2 raise errors when the event times are less than 1e-4. One working solution is to scale the event times (e.g. multiply by 1000) for analysis. TODO: investigate whether we can calculate $X_D(t,x)$ more accurately using the numDeriv package.
- TODO: Extend the generalised survival models to use multiple random effects.
- TODO: Extend the generalised survival models to use automatic differentiation.

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