

Joint modeling of longitudinal and survival data

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Motivation

Joint analysis

New Stata commands for joint analysis

Joint analysis of the PANSS data

Models with more flexible latent associations

Summary

Future work

Acknowledgement

References

- Many studies collect both longitudinal (measurements) data and survival-time data.
- Longitudinal (or panel, or repeated-measures) data are data in which a response variable is measured at different time points such as blood pressure, weight, or test scores measured over time.
- Survival-time or event history data record times until an event of interest such as times until a heart attack or times until death from cancer.

- In the absence of correlation between longitudinal and survival outcomes, each outcome can be analyzed separately.
- Longitudinal analyses include fitting linear mixed models.
- Survival analyses include fitting semiparametric (Cox) proportional hazards models or parametric survival models such as exponential and Weibull.
- When longitudinal and survival outcomes are related, they must be analyzed jointly to avoid potentially biased results.

Joint analyses are useful to:

- Account for informative dropout in the analysis of longitudinal data;
- Study effects of baseline covariates on longitudinal and survival outcomes; or
- Study effects of time-dependent covariates on the survival outcome.

In this presentation, I will concentrate on the first two applications.

- Consider Positive and Negative Symptom Scale (PANSS) data from a clinical trial comparing different drug treatments for schizophrenia (Diggle [1998]).
- We are interested in modeling the total score of the PANSS measurements, which is used to measure psychiatric disorder, over time for each of the drug treatments. The smaller the score the better.
- Six original treatments are combined into three: placebo, haloperidol (reference), and risperidone (novel therapy).
- For details about this study and its analyses, see Diggle (1998) and Henderson (2000).

- We consider a subset of the original data:

```
. use panss
(PANSS scores from a study of drug treatments for schizophrenia)
. describe
Contains data from panss.dta
  obs:           150                PANSS scores from a study of
                                     drug treatments for
                                     schizophrenia
vars:           11                 29 Aug 2016 12:07
size:          3,150              (_dta has notes)
```

variable name	storage type	display format	value label	variable label
id	int	%8.0g		Patient identifier
panss0	int	%8.0g		PANSS score at week 0
panss1	int	%8.0g		PANSS score at week 1
panss2	int	%8.0g		PANSS score at week 2
panss4	int	%8.0g		PANSS score at week 4
panss6	int	%8.0g		PANSS score at week 6
panss8	int	%8.0g		PANSS score at week 8
treat	byte	%11.0g	treatlab	Treatment identifier: 1=Haloperidol, 2=Placebo, 3=Risperidone

nobs	byte	%8.0g		Number of nonmissing measurements, between 1 and 6
droptime	float	%8.0g		Imputed dropout time (weeks)
infdrop	byte	%14.0g	droplab	Dropout indicator: 0=none or noninformative; 1=informative

Sorted by: id

. notes

_dta:

1. Subset of the data from a larger (confidential) randomized clinical trial of drug treatments for schizophrenia
2. Source:
http://www.lancaster.ac.uk/staff/diggle/APTS-data-sets/PANSS_short_data.t
> xt
3. PANSS (Positive and Negative Symptom Scale)

- Listing of a subset of the data:

```
. list id panss* treat if inlist(id,1,2,3,10,19,24,30,42), sepby(nobs) noobs
```

id	panss0	panss1	panss2	panss4	panss6	panss8	treat
1	91	Haloperidol
2	72	Placebo
3	108	110	Haloperidol
10	97	118	Placebo
19	81	71	Risperidone
24	127	98	152	.	.	.	Haloperidol
30	73	74	68	.	.	.	Placebo
42	75	92	117	.	.	.	Risperidone

- Many patients withdrew from the study before completing the measurement schedule—of the 150 subjects, only 68 completed the study.

```
. misstable pattern panss*, freq bypattern
```

```
Missing-value patterns
(1 means complete)
```

Frequency	Pattern				
	1	2	3	4	5
68	1	1	1	1	1
16	1	1	1	1	0
24	1	1	1	0	0
19	1	1	0	0	0
21	1	0	0	0	0
2	0	0	0	0	0
150					

Variables are (1) panss1 (2) panss2 (3) panss4 (4) panss6 (5) panss8

- Over 40% of subjects specified the reason for dropout as “inadequate for response”, which suggests that the dropout may be informative.

```
. tabulate infdrop
```

Dropout indicator	Freq.	Percent	Cum.
None, noninf.	87	58.00	58.00
Informative	63	42.00	100.00
Total	150	100.00	

- Let's first perform standard longitudinal analysis assuming noninformative or random dropout.

```
. use panss_long
(PANSS scores from a study of drug treatments for schizophrenia)
. describe
Contains data from panss_long.dta
  obs:          900                PANSS scores from a study of
                                   drug treatments for
                                   schizophrenia
vars:           6                  29 Aug 2016 12:07
size:          9,900              (_dta has notes)
```

variable name	storage type	display format	value label	variable label
id	int	%8.0g		Patient identifier
week	byte	%9.0g		Time (weeks)
panss	int	%8.0g		PANSS
treat	byte	%11.0g	treatlab	Treatment identifier: 1=Haloperidol, 2=Placebo, 3=Risperidone
nobs	byte	%8.0g		Number of nonmissing measurements, between 1 and 6
panss_mean	float	%9.0g		Observed means over time and treatment

Sorted by: id week

```
. list id week panss treat in 1/16, sepby(id)
```

	id	week	panss	treat
1.	1	0	91	Haloper.
2.	1	1	.	Haloper.
3.	1	2	.	Haloper.
4.	1	4	.	Haloper.
5.	1	6	.	Haloper.
6.	1	8	.	Haloper.
7.	2	0	72	Placebo
8.	2	1	.	Placebo
9.	2	2	.	Placebo
10.	2	4	.	Placebo
11.	2	6	.	Placebo
12.	2	8	.	Placebo
13.	3	0	108	Haloper.
14.	3	1	110	Haloper.
15.	3	2	.	Haloper.
16.	3	4	.	Haloper.

- Consider the following random-intercept model:

$$\text{panss}_{ij} = \beta^L \mathbf{x}_{ij} + U_i + \epsilon_{ij} \quad (1)$$

with m subjects ($i = 1, 2, \dots, m$) and n_i observations per subject ($j = 1, 2, \dots, n_i$), where $\beta^L \mathbf{x}_{ij}$ represents a saturated model with one coefficient for each treat and week combination.

- U_i 's \sim i.i.d. $N(0, \sigma_u^2)$ are random intercepts which induce dependence within subjects.
- ϵ'_{ij} 's \sim i.i.d. $N(0, \sigma_e^2)$ are error terms.

- We use `xtreg`, `mle` to fit a simple random-intercept model by using maximum likelihood (ML) with fixed effects for each combination of treatment and time:

```
. xtset id
      panel variable:  id (balanced)
. xtreg panss i.treat##i.week, mle nolog
Random-effects ML regression              Number of obs   =       685
Group variable: id                       Number of groups =       150
Random effects u_i ~ Gaussian            Obs per group:
                                          min =           1
                                          avg =          4.6
                                          max =           6
                                          LR chi2(17)    =    105.58
                                          Prob > chi2    =     0.0000
Log likelihood = -2861.58
```

	Coef.	Std. Err.	z	P> z	[95% Conf. Interval]
panss					
treat					
Placebo	-2.00	4.14	-0.48	0.629	-10.11 6.11
Risper.	-2.14	4.14	-0.52	0.605	-10.25 5.97

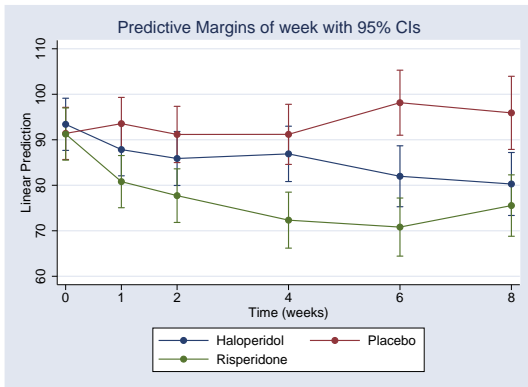
week						
1	-5.55	2.52	-2.21	0.027	-10.49	-0.62
2	-7.51	2.62	-2.87	0.004	-12.64	-2.38
4	-6.50	2.70	-2.40	0.016	-11.80	-1.20
6	-11.42	3.06	-3.73	0.000	-17.41	-5.43
8	-13.12	3.19	-4.12	0.000	-19.36	-6.88
treat#week						
Placebo#1	7.70	3.56	2.16	0.031	0.72	14.68
Placebo#2	7.28	3.80	1.91	0.056	-0.17	14.74
Placebo#4	6.29	4.04	1.56	0.119	-1.63	14.21
Placebo#6	18.17	4.50	4.03	0.000	9.34	26.99
Placebo#8	17.63	4.96	3.56	0.000	7.92	27.35
Risper.#1	-4.91	3.55	-1.38	0.167	-11.86	2.05
Risper.#2	-6.02	3.68	-1.64	0.102	-13.24	1.19
Risper.#4	-12.42	3.85	-3.23	0.001	-19.97	-4.87
Risper.#6	-9.03	4.20	-2.15	0.032	-17.26	-0.79
Risper.#8	-2.60	4.43	-0.59	0.558	-11.29	6.09
_cons	93.40	2.92	31.93	0.000	87.67	99.13
/sigma_u	16.48	1.10			14.47	18.78
/sigma_e	12.49	0.38			11.76	13.26
rho	0.64	0.03			0.57	0.70

LR test of sigma_u=0: chibar2(01) = 353.11

Prob >= chibar2 = 0.000

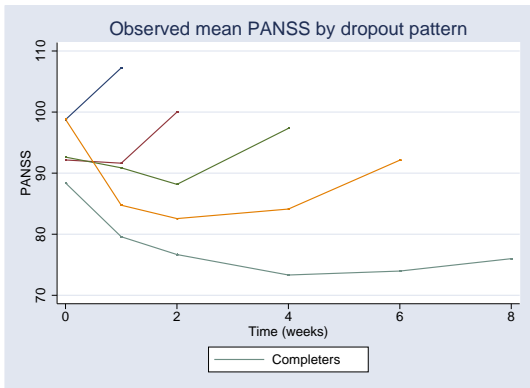
- All three groups demonstrate a decrease in mean PANSS score over time, at least in the first three weeks.

```
. quietly margins i.week, over(treat) predict(xb)
. marginsplot
Variables that uniquely identify margins: week treat
```



- Given that many subjects dropped out of the study because of inadequate response, the observed decrease in PANSS scores may be due to the dropout of subjects with high PANSS scores.
- We can look at the observed mean profiles over time for each missing-value pattern, similarly to Figure 13.4 in Diggle et al. (2002).

```
. keep if nobs>1
(12 observations deleted)
. by week nobs, sort: egen panss_ptrn = mean(panss)
(205 missing values generated)
. qui reshape wide panss_ptrn, i(id week) j(nobs)
. twoway line panss_ptrn* week, sort legend(order(5 "Completers")) ///
> title(Observed mean PANSS by dropout pattern) ytitle(PANSS)
```



- There is a step increase in the mean PANSS score immediately prior to dropout for all dropout patterns except completers.
- This provides strong empirical evidence that dropout is related to PANSS scores and is thus informative (nonrandom).

- We may also be interested in a dropout process itself. For example, is there a difference between dropout rates because of “inadequate response” among groups?
- We can use standard methods of survival analysis to answer this question.
- We can treat dropout time as our analysis time and whether the dropout is because of inadequate response as our event of interest or failure.

- Data description:

```
. use panss_surv
(Dropout times for study of drug treatments for schizophrenia)
. describe
Contains data from panss_surv.dta
  obs:                150                Dropout times for study of drug
                                          treatments for schizophrenia
  vars:                4                 29 Aug 2016 12:07
  size:                1,200            (_dta has notes)
```

variable name	storage type	display format	value label	variable label
id	int	%8.0g		Patient identifier
droptime	float	%8.0g		Imputed dropout time (weeks)
infdrop	byte	%14.0g	droplab	Dropout indicator: 0=none or noninformative; 1=informative
treat	byte	%11.0g	treatlab	Treatment identifier: 1=Haloperidol, 2=Placebo, 3=Risperidone

Sorted by: id

```
. list in 1/10
```

	id	droptime	infdrop	treat
1.	1	.704	None or noninf.	Haloper.
2.	2	.74	None or noninf.	Placebo
3.	3	1.121	Informative	Haloper.
4.	4	1.224	Informative	Haloper.
5.	5	1.303	None or noninf.	Haloper.
6.	6	1.541	Informative	Haloper.
7.	7	1.983	Informative	Haloper.
8.	8	1.035	Informative	Placebo
9.	9	1.039	None or noninf.	Placebo
10.	10	1.116	Informative	Placebo

- Cox proportional hazards model:

$$h_i(t|\text{treat}) = h_0(t) \exp(\beta_1^S 1.\text{treat}_i + \beta_2^S 2.\text{treat}_i + \beta_3^S 3.\text{treat}_i) \quad (2)$$

where t is the dropout time `droptime` and $i = 1, 2, \dots, m$.

- Baseline hazard $h_0(t)$ is left unspecified.
- A constant term β_0^S is absorbed into the baseline hazard.
- Coefficients β_1^S , β_2^S , and β_3^S model subject-specific hazards as a function of the treatment group. In general, covariates may also depend on time t .
- Subject-specific hazards are proportional.
- Exponentiated coefficients are hazard ratios.

- Declare survival-time data:

```
. stset droptime, failure(infdrop)
      failure event:  infdrop != 0 & infdrop < .
obs. time interval:  (0, droptime]
exit on or before:  failure
```

```
      150 total observations
       0 exclusions
```

```
      150 observations remaining, representing
       63 failures in single-record/single-failure data
863.624 total analysis time at risk and under observation
              at risk from t =           0
          earliest observed entry t =       0
              last observed exit t =       8.002
```


- Fit Cox model:

```
. stcox i.treat
      failure _d:  infdrop
      analysis time _t:  droptime
Iteration 0:  log likelihood = -293.97982
Iteration 1:  log likelihood = -288.97387
Iteration 2:  log likelihood = -288.86504
Iteration 3:  log likelihood = -288.86498
Refining estimates:
Iteration 0:  log likelihood = -288.86498
Cox regression -- Breslow method for ties
No. of subjects =          150          Number of obs   =          150
No. of failures =           63
Time at risk    = 863.6239911
Log likelihood   = -288.86498
LR chi2(2)      =          10.23
Prob > chi2     =          0.0060
```

_t	Haz. Ratio	Std. Err.	z	P> z	[95% Conf. Interval]
treat					
Placebo	1.81	0.53	2.04	0.041	1.02 3.21
Risper.	0.68	0.24	-1.12	0.262	0.34 1.34

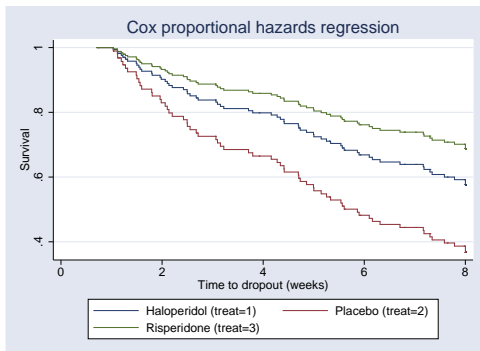
- Redisplay results as coefficient estimates (for later comparison):

```
. stcox, nohr
Cox regression -- Breslow method for ties
No. of subjects =          150          Number of obs   =          150
No. of failures =           63
Time at risk    = 863.6239911
Log likelihood  = -288.86498
LR chi2(2)     =          10.23
Prob > chi2    =          0.0060
```

_t	Coef.	Std. Err.	z	P> z	[95% Conf. Interval]	
treat						
Haloper.	0.00	(empty)				
Placebo	0.59	0.29	2.04	0.041	0.02	1.16
Risper.	-0.39	0.35	-1.12	0.262	-1.07	0.29

- Plot survivor functions in three treatment groups:

```
. stcurve, survival at1(treat=1) at2(treat=2) at3(treat=3)
```



- The placebo group has the highest dropout rate due to inadequate response whereas the risperidone group has the lowest dropout rate.
- But dropout rates also depend on PANSS scores.

- Whether we are interested:
 - In the longitudinal analysis of PANSS trajectory over time in different groups,
 - In the survival analysis comparing dropout rates among the groups, or
 - In both types of analysis,we cannot perform them separately, given that the two outcomes may be correlated.
- We should consider joint analysis of these data.

- Joint analysis should be able to incorporate the specific features of longitudinal and survival data.
- Joint analysis should be equivalent to the corresponding separate analysis in the absence of an association between the longitudinal and survival outcomes.
- Tsiatis et al. (1995), Wulfsohn and Tsiatis (1997), and Henderson et al. (2000) considered a joint model that links the longitudinal and survival outcomes through a shared latent process.

- Let's fit a model that accounts for informative dropout.
- Consider the following joint random-intercept Cox model based on separate models (1) and (2):

$$\begin{aligned}\text{panss}_{ij} &= \beta^L \mathbf{x}_{ij} + U_i + \epsilon_{ij} \\ h_i(t) &= h_0(t) \exp(\beta^S \text{i.treat}_i + \gamma U_i)\end{aligned}\quad (3)$$

- Random intercepts U_i 's are now shared between the two models and induce dependence between the longitudinal outcome `panss` and survival outcome `droptime`.
- More generally, I will refer to model (3) as a joint random-intercept Cox model, in which survival outcome is modeled semiparametrically using the Cox model.

- You can use forthcoming, user-written suite `jm` to perform joint analysis of longitudinal and survival data.
- Command `jmxstset` declares your longitudinal and survival data.
- Command `jmxstcox` fits joint random-intercept Cox models, similar to model (3).
- Command `jmxstcurve` plots survivor, hazard, and cumulative hazard functions after `jmxstcox`.
- Other Stata postestimation features such as `predict`, `test`, `nlcom`, `margins`, etc. are also available.

- To fit joint models using `jmxststcox`, you must first declare your longitudinal and survival data using `jmxststset`.
- Longitudinal and survival data are typically saved in different files. To perform estimation, all data should be in one file with longitudinal data saved in a long format (with multiple observations per subject saved in rows).
- `jmxststset` provides a syntax that combines the two datasets and performs declaration, and provides a syntax that declares an already combined dataset.

- `jmxxtstset` combines the syntaxes of `stset` and `xtset`.
- Syntax for the combined dataset:

```
. jmxxtstset idvar timevar, xt(is_xt)|st(is_st) failure(failvar) [stsetopts]
```

is_xt and *is_st* are binary variables identifying longitudinal and survival observations, respectively; only one of them must be specified in the respective option.

- Syntax for separate datasets with survival dataset in memory:

```
. use survfile  
. jmxxtstset idvar timevar using longfile, st failure(failvar) [stsetopts]
```

- Syntax for separate datasets with longitudinal dataset in memory:

```
. use longfile  
. jmxxtstset idvar timevar using survfile, xt failure(failvar) [stsetopts]
```

- Command `jmxststcox` performs estimation.
- It fits a random-intercept Cox model to the survival and longitudinal outcomes.
- `jmxststcox` uses nonparametric ML to estimate model parameters. The estimation method is an expectation-maximization algorithm. The standard errors are obtained using the observed information matrix (Louis 1982).

- Command `gsem` (`help gsem`) can be used to fit joint models with flexible specification of latent processes, but in which survival outcome is modeled parametrically.
- User-written command `stjm` (Crowther et al. 2013) can be used to fit joint random-intercept and random-coefficient models. The survival outcome is again modeled parametrically, but flexible parametric survival models (Royston and Lambert 2011) are also supported.
- User-written command `jmxtstcox` currently supports only joint random-intercept models, but it allows to model the survival outcome semiparametrically, without any parametric assumptions for the baseline hazard.

- Let's now analyze PANSS scores and dropout times jointly by fitting the random-intercept Cox model (3).
- The longitudinal data are saved in `panss_long.dta` and the survival data are saved in `panss_surv.dta`.
- We first use `jmxtstset` to combine survival and longitudinal datasets and to declare the combined data:

```
. use panss_surv
(Dropout times for study of drug treatments for schizophrenia)
. jmxtstset id droptime using panss_long, st failure(infdrop)
```

```
-----LONGITUDINAL-----
           id:  id
      filename: panss_long.dta
```

```
      900  total observations
         0  exclusions
```

```
      900  observations remaining
      150  subjects
```

```

-----SURVIVAL-----
      id: id
      failure event: infdrop != 0 & infdrop < .
obs. time interval: (droptime[_n-1], droptime]
      exit on or before: failure
-----
      150 total observations
       0 exclusions
-----
      150 observations remaining, representing
      150 subjects
       63 failures in single-failure-per-subject data
      863.624 total analysis time at risk and under observation
                                at risk from t =           0
      earliest observed entry t =           0
                                last observed exit t =      8.002

```

- We now use `jmxtstcox` to fit the joint model:

```
. jmxtstcox (_xt: panss i.treat##i.week) (_st: i.treat), nolog
      longitudinal depvar:  panss
                        failure _d:  infdrop
                        analysis time _t:  droptime
Joint model of longitudinal and survival data
Breslow method for ties
Subject id: id
Longitudinal (_xt):
No. of subjects = 150
No. of obs      = 685
Total subjects = 150
Survival (_st):
No. of subjects = 150
No. of obs      = 150
No. of failures = 63
Time at risk    = 863.62
Wald chi2(19)   = 112.90
Prob > chi2     = 0.0000
```

	Coef.	Std. Err.	z	P> z	[95% Conf. Interval]	
panss						
treat						
Placebo	-2.00	4.16	-0.48	0.631	-10.16	6.16
Risper.	-2.14	4.16	-0.51	0.607	-10.30	6.02
week						
1	-5.55	2.51	-2.21	0.027	-10.48	-0.63
2	-7.24	2.61	-2.77	0.006	-12.36	-2.13
4	-6.12	2.70	-2.27	0.023	-11.40	-0.83
6	-10.61	3.05	-3.48	0.001	-16.59	-4.63
8	-12.20	3.18	-3.84	0.000	-18.43	-5.97
treat#week						
Placebo#1	7.69	3.55	2.17	0.030	0.73	14.66
Placebo#2	7.65	3.79	2.02	0.044	0.22	15.09
Placebo#4	7.03	4.03	1.75	0.081	-0.86	14.93
Placebo#6	18.74	4.49	4.18	0.000	9.95	27.54
Placebo#8	18.43	4.94	3.73	0.000	8.75	28.11
Risper.#1	-4.91	3.54	-1.39	0.166	-11.84	2.03
Risper.#2	-6.08	3.67	-1.65	0.098	-13.28	1.13
Risper.#4	-12.30	3.84	-3.20	0.001	-19.83	-4.77
Risper.#6	-9.12	4.19	-2.18	0.029	-17.33	-0.91
Risper.#8	-2.82	4.42	-0.64	0.524	-11.48	5.85
_cons	93.40	2.93	31.85	0.000	87.65	99.15

<code>_t</code>							
treat							
Placebo	0.77	0.34	2.23	0.026	0.09	1.44	
Risper.	-0.49	0.39	-1.26	0.207	-1.26	0.27	
/ gamma	0.05	0.01			0.04	0.07	
/ sigma2_u	281.22	37.30			208.11	354.34	
/ sigma2_e	155.29	9.47			136.73	173.85	

LR test of gamma = 0: chi2(1) = 37.41

Prob >= chi2 = 0.0000

- The association parameter γ has an estimate of 0.05 with a 95% CI of (0.04, 0.07), which implies a positive association between PANSS scores and dropout times—the higher the PANSS score, the higher the chance of dropout.
- The LR test of no latent association ($H_0: \gamma = 0$) with $\chi_1^2 = 37.41$ provides strong evidence against a random-dropout model.

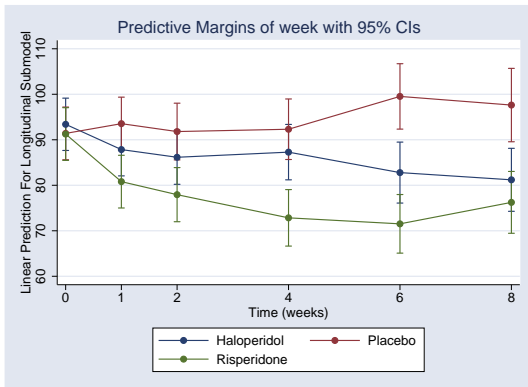
- The estimated random-intercept variance is slightly larger under the joint, informative dropout model.

Variable	inform	noninf
sigma2_u		
_cons	281.22	271.75
	37.30	36.19
	0.00	0.00
sigma2_e		
_cons	155.29	155.95
	9.47	9.55
	0.00	0.00

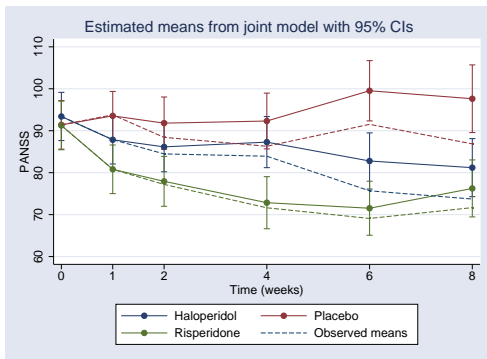
legend: b/se/p

- As with `xtreg`, we can compute and plot estimated mean PANSS profiles after `jmxststcox`.

```
. qui margins i.week, over(treat) predict(xb xt)
. marginsplot
Variables that uniquely identify margins: week treat
```



- We can overlay the estimated mean profiles with the observed mean profiles.



- The estimated mean profiles from the joint model are higher than the observed mean profiles because the former represent “dropout-free” profiles—subjects with high PANSS scores tend to drop out, which leads to lower observed mean values.

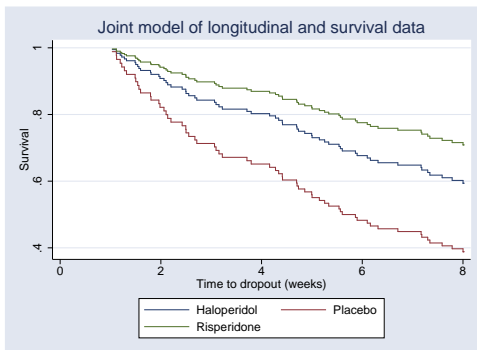
- We can compare estimates from joint and separate Cox models:

Variable	joint	stcox
treat		
Haloper.	(base)	(base)
Placebo	0.77	0.59
	0.34	0.29
	0.03	0.04
Risper.	-0.49	-0.39
	0.39	0.35
	0.21	0.26

legend: b/se/p

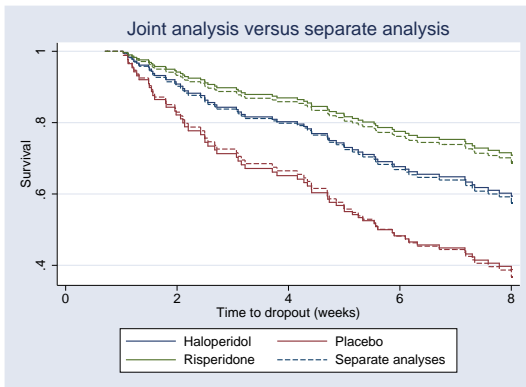
- We can plot marginal survivor functions of times to dropout in each group.

```
. jmxstcurve, survival at1(treat=1) at2(treat=2) at3(treat=3)
```



- As with separate analysis, the placebo group has the highest “informative” dropout rate whereas the risperidone group has the lowest dropout rate.

- In fact, survival estimates from joint and separate analyses are similar:



- Random-intercept model (3) can be extended to allow for more flexible latent associations motivated by practice; see Henderson (2000) for details.
- For example, a joint random-coefficient Cox model additionally includes a random slope on time in the longitudinal model and an association through the random slope in the survival model.

$$\begin{aligned} \text{panss}_{ij} &= \beta^L \mathbf{x}_{ij} + U_{1i} + \text{week} \times U_{2i} + \epsilon_{ij} \\ h_i(t) &= h_0(t) \exp(\beta^S \text{i.treat}_i + \gamma_1 U_{1i} + \gamma_2 U_{2i}) \quad (4) \end{aligned}$$

- A joint random-trajectory Cox model extends the random-coefficient model (4) to include an entire stochastic longitudinal trajectory.

$$\begin{aligned} \text{panss}_{ij} &= \beta^L \mathbf{x}_{ij} + U_{1i} + \text{week} \times U_{2i} + \epsilon_{ij} \\ h_i(t) &= h_0(t) \exp(\beta^S \text{i.treat}_i + \gamma_1 U_{1i} + \gamma_2 U_{2i} + \gamma_3 W_i(t)) \\ W_i(t) &= U_{1i} + t \times U_{2i} \quad (5) \end{aligned}$$

- Semiparametric Cox submodels in (3), (4), and (5) can be replaced with a parametric survival model, if appropriate.
- For example, with an exponential model:

$$h_i(t) = t \exp(\beta^S \text{i.treat}_i + \gamma U_i) \quad (3a)$$

- Or, with a Weibull model:

$$h_i(t) = pt^{p-1} \exp(\beta^S \text{i.treat}_i + \gamma U_i) \quad (3b)$$

- Such parametric models can be fit using, for example, `gsem`, but software for the corresponding semiparametric models is not available yet.

- For example, a joint random-intercept model using gsem:

```
. gsem (panss <- i.treat##i.week U[id]@1)
> (droptime <- i.treat U[id]@gamma, family(weibull, failure(infdrop))
```

- A joint random-coefficient model:

```
. gsem (panss <- i.treat##i.week U1[id]@1 c.week#U2[id]@1)
> (droptime <- i.treat U1[id]@gamma1 U2[id]@gamma2,
>      family(weibull, failure(infdrop))),
> covstructure(U1[id] U2[id], unstructured)
```

Summary

- Joint analysis of longitudinal and survival outcomes is necessary to obtain unbiased inference when the two outcomes are correlated.
- Joint analysis can be used, for example, 1) to evaluate effects of baseline covariates on longitudinal and survival outcomes, 2) to evaluate effects of time-dependent covariates on survival outcome; and 3) to account for informative dropout in longitudinal analysis.
- You can use user-written command `jmxststcox` to fit a joint random-intercept Cox model.
- You can use `gsem` to fit joint models that can accommodate more flexible specifications of a latent process and noncontinuous longitudinal outcomes. The survival outcome, however, is modeled parametrically.
- Also see user-written command `stjm` for fitting flexible parametric joint models of longitudinal and survival data.

Future work

- Support of semiparametric Cox models with more flexible latent associations such as a random-coefficient model (4) and a random-trajectory model (5).
- Support of noncontinuous longitudinal outcomes including binary and count outcomes.
- Support of nonproportional hazards via transformation survival models (Zeng and Lin 2007).
- More postestimation features such as dynamic predictions and model diagnostics for joint analysis of longitudinal and survival data.

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