

Discussion: Modelling baseline excess hazard function in relative survival

Patrick Royston, MRC Clinical Trials Unit, London

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Statement of Interest!

- I am biased!
- Nelson's approach is essentially that of Royston & Parmar (2002)
- 5 years later, I still like the approach
- I therefore encourage Nelson and others to research this family of models further

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Bernard Rachet

- Time and covariates are not (or need not be) categorised
- FPs and/or regression splines as functions of time for modelling baseline excess hazard
- ... also for interactions of covariates with time
- Principled approach to model selection using MFP

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Issues with Rachet's approach

- Modelling the excess hazard with FPs/splines may give artefacts in the curves
- "How flexible?" seems to be a general issue
 - Particularly with large datasets
 - Criteria for model selection?
- Selecting interaction models – may be too flexible, leading to instability/possible interpretation problems
 - Breast cancer example
- May need expert "tuning" to get good results

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Chris Nelson

- Main difference: modelling *cumulative* excess hazard function
- Splines in log t for this function
- Can also model covariates in similar fashion

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Some issues with categorisation (Nelson)

- Several methods split the timescale into manageable chunks for estimation purposes
 - Necessary with large datasets
- Collapse on time and grouped covariates
 - Can't handle continuous covariates
- Piecewise models could be considered as biologically implausible, both for time and covariates (also, biased?)

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- Many parameters for time-dependent effects
 - Time chunks x covariate groups = lots
 - Overfitted, unrealistic
 - Can lead to problems in smaller datasets, e.g. zero cells
- If have many observations, may get computation-time problems on split data
- Methods that do *not* split time require integration to estimate survival function
 - Because estimating excess hazard, not cumulative

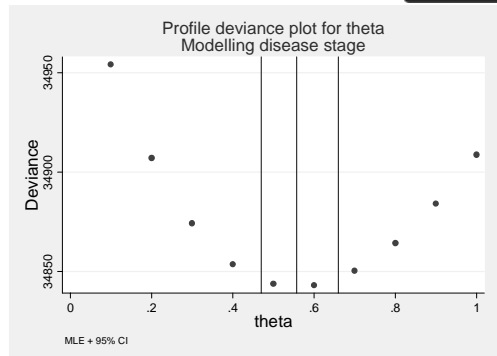
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Comments

- For me, Nelson’s approach is close to the method of choice
- Could substitute fractional polynomials for splines in some applications (Lambert, Rachet)
 - Model baseline cumulative excess hazard
 - Model continuous covariates
- Don’t forget the other link functions
 - $\Theta = 1$ (cumulative proportional odds)
 - $\Theta \neq 0, 1$ (can be estimated from the data)

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Colon cancer data – stage known



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Regression splines – in favour

- Parametric
 - Concise description of function available
- Flexible
 - Can cope with many different curve shapes
- Can control flexibility via number of knots
 - Not usually too sensitive to exact knot positioning
- Easily incorporated in regression models
 - No special computational methods required

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Regression splines – against

- Baseline cumulative excess hazard function not guaranteed monotonic in time
- Model (knot) selection – is AIC satisfactory?
 - No simulation evidence available
- Can get artefacts in fitted function
 - Problems with over-interpretation
 - May not be transportable to other datasets (validation)
- Highly correlated terms
 - Need to orthogonalise

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A difficult issue: model selection

- Scale of model
 - Hazard, odds, probit, etc.
- Baseline (spline terms)
 - Done separate or simultaneously with covariates?
- How to do scale, variable and function selection?
 - E.g. when have prognostic factors in cancer
- Many possible covariate x time interactions give serious multiplicity problem
 - Forward selection?
- Assessing model stability (e.g. bootstrap)

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Paul Lambert

- Cure models – mixture and non-mixture formulations
- Seems particularly appropriate approach in relative survival context
 - In most primary cancers, some patients will be cured
 - For metastatic (e.g. stage 4) cancer, cure proportion is nearly 0
- Preference for non-mixture model
 - More realistic
 - Is also interpretable as a mixture model

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Issues

- Are the models identifiable?
 - Model with excess hazards converging towards zero may fit equally well – and could use more flexible distributional specification
- Robustness - estimate of cure fraction (π) depends on choice of parametric survival distribution
- Is it possible to fit Royston-Parmar-Nelson distributions to $S(t)$?
- How to choose a model – many possibilities?

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Ula Nur

- Handle missing covariate values by multiple imputation (MI)
- Create m complete datasets (e.g. $m = 10$)
- Requires a (complex) imputation model
 - To make MAR assumption more plausible
- Do MI using `ice`
- Combine results from models fit by `strs`, `strel` etc using `micombine` or `mim`
 - `mim` is preferred command now

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Issues

- Handling of categorical covariates needs care
 - Use `passive()` and `substitute()` options in `ice`
- Not clear how/whether survival time should be transformed in `ice` model
 - General issue in survival analysis with missing data
 - Probably depends on substantive model to be fitted
- Since covariate x time interactions are a feature of RS models, should consider including them in `ice` model
 - Omission will cause bias towards zero

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Future

- What are the main aims of relative survival models?
- If *description* then *goodness of fit* and *interpretability* are important
- If *prediction* in independent data then *external validation* is important
- What implications would these aims have for the preferred method of modelling?
- E.g. (when) should cure models be the approach of choice?

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Further applications

- Meta analysis of relative survival functions across hospitals, regions, countries, etc?
- Role in regular prognostic modelling?
- Role in economic evaluation of cancer therapies using data from clinical trials?
 - Estimate disease-related years of life lost

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Stata

- Standardise popmort file specification for use by strel, strsm* and other applications?
- Recommendation of particular approaches/Stata commands to answer specific questions?