Semiparametric Regression for Dual Population Mortality

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Abstract:
Parameter shrinkage applied optimally can always reduce error and projection variances from those of maximum likelihood estimation. Many variables that actuaries use are on numerical scales, like age or year, which require parameters at each point. Rather than shrinking these towards zero, nearby parameters are better shrunk towards each other. Semiparametric regression is a statistical discipline for building curves across parameter classes using shrinkage methodology. It is similar to but more parsimonious than cubic splines. We introduce it in the context of Bayesian shrinkage and apply it to joint mortality modeling for related populations, with Swedish and Danish mortality as an illustration. Bayesian shrinkage of slope changes of linear splines is an approach to semiparametric modeling that evolved in the actuarial literature. It has some theoretical and practical advantages, like closed-form curves, direct and transparent determination of degree of shrinkage and of placing knots for the splines, and quantifying goodness of fit. It is also relatively easy to apply to the many nonlinear models that arise in actuarial work.

Keywords: Semiparametric regression, Joint mortality, Parameter shrinkage, Bayesian shrinkage, MCMC
1 Introduction
Actuaries often need to estimate mortality models for specialized subpopulations of a larger population, e.g., for pricing products for high-net-worth individuals. Dual mortality modeling seeks to incorporate the data of the larger population while allowing the subpopulation model to differ to the extent it has reliability. We develop a method to apply Bayesian shrinkage at this step. This lets the parameters of the two models differ to the extent that this improves the joint posterior distribution.

The method explored is to use Bayesian semiparametric models for both populations but make the model for the smaller population the larger-population model plus a semiparametric model for the differences, with both fit simultaneously. We build up increasingly complex models, culminating in a negative binomial version of the Hunt-Blake model (Hunt and Blake (2014)). As an illustration, we apply this to jointly model male mortality for Denmark and Sweden. This is not a typical dual-mortality modeling application, as Denmark with 5.6 million people would often be modeled on its own, even though Sweden is almost twice as populous. However the data is publicly available and it lets us detail the steps of the process.

Quite a few actuarial papers have addressed joint age-period-cohort (APC) modeling of related datasets in reserving and mortality. Li and Lee (2005) introduce the concept of using a joint stochastic age-period process for mortality modeling of joint populations, with mean reverting variations from the joint process for each population. A similar approach by Jarner and Kryger (2009) models a large and a small population with the small population having a multi-factor mean-zero mean reverting spread in log mortality rates. Dowd et al. (2011) model two populations that can be of comparable or different sizes with mean reverting stochastic spreads for both period and cohort trends. A period-trend spread in an APC model is estimated by Cairns et al. (2011), who use Bayesian Markov-Chain Monte Carlo (MCMC) estimation. They assume fairly wide priors, not shrinkage priors. This uses the capability of MCMC to estimate difficult models. Antonio, Bardoutsos, and Ouburg (2015) estimate the model of Li and Lee (2005) by MCMC.

For some time actuaries have used precursors to semiparametric regression, like linear and cubic spline curves fit across parameter types. Semiparametric regression uses parameter shrinkage methodology to produce an optimized degree of parsimony in the curve construction. The standard building block for this now is O’Sullivan penalized cubic splines (O’Sullivan (1986)). Harezlak, Ruppert, and Wand (2018) provides good background, with R code. Here we use penalized linear splines, which have some advantages for the models we study.

The actuarial use of shrinkage for linear-spline models started with Barnett and Zehnwirth
(2000), who applied it to APC models for loss reserving, using an ad hoc shrinkage methodology. Venter, Gutkovich, and Gao (2019) use frequentist random-effects methods for linear-spline shrinkage for reserving and mortality models, including joint modeling of related loss triangles. Venter and Şahin (2018) fit the Hunt-Blake mortality model with Bayesian shrinkage of linear splines, determining the shrinkage level by cross-validation. Gao and Meng (2018) use Bayesian shrinkage on cubic splines for loss reserving models, getting the shrinkage level by a fully-Bayesian method. Here we use linear splines on a Hunt-Blake model with a fully-Bayesian estimation approach that combines lasso with MCMC to speed convergence.

Shrinkage methods are discussed in Section 2 and Section 3 has the application to actuarial models. Section 4 works through the steps for fitting the joint model, and Section 5 concludes.

2 Shrinkage Methodologies

The use of shrinkage-type estimation to improve fitting and projection accuracy traces back to actuarial credibility theory, then to various coefficient-shrinkage methods for regression, which we work though in this section. These have gotten easier to apply over time, and also have been extended to building curves across levels of some variables.

2a Credibility

Shrinking estimates towards the grand mean started with combined estimation of a number of separate means. A typical example is batting averages for a group of baseball players. Stein (1956) showed that properly shrinking such means towards the overall mean always reduces estimation and prediction variances from those of maximum likelihood estimation (MLE) when there are three or more means being estimated. Similar methods have been part of actuarial credibility theory since Mowbray (1914), and in particular Bühlmann (1967). According to the Gauss-Markov Theorem, MLE is the minimum-variance unbiased estimate. Credibility biases the individual estimates towards the grand mean, but reduces the estimation errors. Thus having more accurate estimates comes at the cost of possibly asymmetric even though smaller confidence intervals. This holds in all of the shrinkage methods below.

2b Regularization

For regression models, Hoerl and Kennard (1970), showed that some degree of shrinking coefficients towards zero always produces lower fitting and prediction errors than MLE. This is related to credibility, as in their ridge-regression approach, variables are typically scaled to have mean zero, variance one, so reducing coefficients shrinks the fitted means towards the constant term, which is the overall mean. Their initial purpose was to handle regression with correlated independent variables, and they applied a more general procedure for ill-posed problems known as Tikhonov regularization, from Tikhonov (1943). As a result, shrinkage
methods are also referred to as regularization.

With coefficients $\beta_j$, ridge regression minimizes the negative loglikelihood (NLL) plus a selected shrinkage factor $\lambda$ times the sum of squares of the coefficients. Thus it minimizes:

$$NLL + \lambda \sum_j \beta_j^2$$

The result of Hoerl and Kennard (1970) is that there is always some $\lambda > 0$ for which ridge regression gives a lower estimation variance than does MLE. They did not have a formula to optimize $\lambda$, but could find reasonably good values by cross validation – that is, by testing prediction on subsamples omitted from the fitting.

In the 1990s, lasso, which minimizes:

$$NLL + \lambda \sum_j |\beta_j|$$

became a popular alternative. It shrinks some coefficients exactly to zero, so it provides variable selection as well as shrinkage estimation. A modeler can start off with a long list of variables and use lasso to find combinations that work well together with parameter shrinkage.

**2c Random Effects**

Random effects modeling provides a more general way to produce such shrinkage. Historically it postulated mean-zero normal distributions for the $\beta_j$, with variance parameters to be estimated, and optimized the joint likelihood, which is the product of those normal densities with the likelihood function. This also pushes the coefficients towards zero since they start out having mean zero, mode zero distributions. This can have a different variance for each $\beta_j$, which can also be correlated. Ridge regression arises from the special case where there is a single variance assumed for all the coefficients. Using the normal distribution for this shrinkage is common but it is not a requirement. If the double exponential (Laplace) distribution is assumed instead, lasso becomes a special case as well.

We start by assuming the random effects have independent mean-zero normal distributions with a common variance. Given a vector $b$ of random effects and a vector $\beta$ of parameters (fixed effects), with respective design matrices $Z$ and $X$, and a vector of observations $y$, the model is:

$$y = X\beta + Zb + \epsilon$$

$$b \sim N(0, \theta \sigma^2)$$
\[ \epsilon \sim N(0, \sigma^2) \]

Sometimes fitting the model is described as estimating the parameters and projecting the random effects. A popular way to fit the model is to maximize the joint likelihood, which is:

\[ p(b, y; \beta) = p(y|\beta, b)p(b) \]

This is the likelihood times the probability of the random effects. The mean-zero distribution for the bs pushes their projections towards zero, so this is a form of shrinkage. Matrix estimation formulas have been worked out for this similar to those for regression.

Now we consider instead the double exponential, or Laplace, distribution for the random effects. This is an exponential distribution for positive values of the variable, and is exponential in \(-b\) for negative values. Its density is

\[ p(b|\lambda) = 0.5\lambda e^{-\lambda|b|} \]

Suppose all the random effects have the same \(\lambda\). Then the negative joint loglikelihood is:

\[ -\log[p(b, y; \beta)] = \log(2) - \log(\lambda) + \sum_j \lambda|b_j| + NLL \]

The constants do not affect the minimization, so for a fixed given value of \(\lambda\), this is the lasso estimation. If \(\lambda\) is considered a parameter to estimate, then the \(\log(\lambda)\) term would be included in the minimization, giving an estimate of \(\lambda\) as well. This same calculation starting with a normal prior gives ridge regression. Thus these are both special cases of random effects, and both provide estimates of the shrinkage parameter.

Lasso and ridge regression have problems with measures of penalized NLL like AIC and BIC. These rely on parameter counts, but shrunk parameters do not use as many degrees of freedom, because they have less ability to influence fitted values. Because of this, cross validation is popular for model selection for frequentist shrinkage. There is usually some degree of subjectivity involved in the choice of cross-validation approach. We will see below that Bayesian shrinkage has a fast automated cross-validation method for penalizing the NLL. It provides a good estimate of what the likelihood would be from a new sample from the same population, which is the goal of AIC as well.

**2d Bayesian Shrinkage**

Bayesian shrinkage is quite similar to random effects, with normal or Laplace priors used for the parameters like they are used for postulated distributions of the effects. Ridge regression
and lasso estimates are produced as the posterior modes. The Bayesian and frequentist
approaches are closer than in the past, as random effects are similar to prior distributions
and the Bayesian prior distributions are not necessarily related to prior beliefs. Rather they
are some of the postulated distributions assumed in the model, just like random effects are,
and they can be rejected or modified based on the posteriors that are produced.

In the Bayesian approach, a parameter might be assumed to be normally distributed with
mean zero, or Laplace or otherwise distributed. For a sample \( X \) and parameter vector \( \beta \),
with prior \( p(\beta) \) and likelihood \( p(X|\beta) \), the posterior is given by Bayes Theorem as:

\[
p(\beta|X) = \frac{p(X|\beta)p(\beta)}{p(X)}
\]

Here the probability of the data, \( p(X) \), is not usually known but it is a constant. Since
\( p(\beta|X) \) must integrate to 1, the posterior is determined by \( p(X|\beta)p(\beta) \).

Note that if \( \beta \) were random effects, the numerator of \( p(\beta|X) \) would be the joint likelihood.
Thus maximizing the joint likelihood gives the same estimate as the mode of the posterior
distribution. The main difference between Bayesians and frequentists now is that the former
mostly use the posterior mean, and the latter use the posterior mode. Bayesians tend to
suspect that the mode could be overly responsive to peculiarities of the sample. Frequentists
come from a lifetime of optimizing goodness of fit of some sort, which the mode does.

Bayesians generate a sample of the posterior numerically using MCMC. This ranks parameter
samples using the joint likelihood, since this is proportional to the posterior. MCMC has
a variety of sampling methods it can use. The original one was the Metropolis algorithm,
which has a generator for the possible next sample using the latest accepted sample. If the
new sample parameters have a higher joint probability, it is accepted. If not, a random draw
is used to determine whether it is accepted or not. An initial group of samples is considered
to be warmup and is eventually eliminated, and the remainder using this procedure have
been shown to follow the posterior distribution.

We feel that random effects estimation can also make use of MCMC to calculate the posterior
mean, but under a different framework. Now we will just consider the case where there are
no fixed effects to estimate – all the effects in the model are random effects. A distribution is
postulated for the effects, including a postulated distribution for \( \lambda \). We know how to calculate
the mode of the conditional distribution of the effects given the data, namely by maximizing
the joint likelihood. By the definition of conditional distributions, the joint likelihood =

\[
p(X|\beta)p(\beta) = p(X, \beta) = p(X)p(\beta|X)
\]
Thus the conditional distribution of the effects given the data is proportional to the joint likelihood, and so can be estimated by MCMC.

What the Bayesians would call the posterior mean and mode of the parameters can instead be viewed as the conditional mean and mode of the effects given the data. With MCMC, the effects can be projected by the conditional mean instead of the conditional mode that was produced by maximizing the joint likelihood. This does not require Bayes Theorem, subjective prior and posterior probabilities, or distributions of parameters, so removes many of the objections frequentists might have to using MCMC. Admittedly Bayes Theorem is obtained by simply dividing the definition of conditional distribution by \( p(X) \), but you do not have to do that division to show that the conditional distribution of the effects given the data is proportional to the joint likelihood.

2e Goodness of fit from MCMC

One advantage of using MCMC, whether the mean or mode is used, is that there is a convenient goodness-of-fit measure. The leave-one-out, or loo, cross validation method is to refit the model once for every sample point, leaving out that point in the estimation. Then the likelihood of that point is computed from the parameters fit without it. The cross-validation measure, the sum of the loglikelihoods of the omitted points, has been shown to be a good estimate of the loglikelihood for a completely new sample from the same population, so it eliminates the sample bias in the NLL calculation. This is what AIC, etc. aim to do as well.

But doing all those refits can be very resource intensive. Gelfand (1996) developed an approximation for an omitted point’s likelihood from a single MCMC sample generated from all the points. He used the numerical integration technique of importance sampling to estimate a left-out point’s likelihood by the weighted average likelihood across all the samples, with the weight for a sample proportional to the reciprocal of the point’s likelihood under that sample. That gives greater weight to the samples that fit that point poorly, and turns out to be a good estimate of the likelihood that it would have if it had been left out of the estimation. This estimate for the likelihood of the point turns out to be the harmonic mean over the samples of the point’s probability in each sample. With this, the MCMC sample of the posterior distribution is enough to calculate the loo goodness-of-fit measure.

This gives good but volatile estimates of the loo loglikelihood. Vehtari, Gelman, and Gabry (2017) addressed that by a method similar to extreme value theory – they fit a Pareto to the probability reciprocals and use the Pareto values instead of the actuals for the largest 20% of the reciprocals. They call this “Pareto-smoothed importance sampling.” It has been extensively tested and is becoming widely adopted. Their penalized likelihood measure is
labeled $\hat{elpd}_{\text{loo}}$, standing for “expected log pointwise predictive density.” Here we call $-\hat{elpd}_{\text{loo}}$ simply loo. It is the NLL plus a penalty, so is a penalized likelihood measure.

The fact that this is a good estimate of the NLL without sample bias comes with a caveat. The derivation of that assumes that the sample comes from the modeled process. That is a standard assumption but in financial areas, models are often viewed as approximations of more complex processes. Thus a new sample might not come from the process as modeled. Practitioners sometimes respond to this by using slightly under-fit models – that is more parsimonious models with a bit worse fit than the measure finds optimal.

Using cross validation to determine the degree of shrinkage usually requires doing the estimation for each left-out subsample for trial values of the shrinkage constant $\lambda$. With Bayesian shrinkage you still have to compute loo for various $\lambda$s, but not by omitting points, as loo already takes care of that. This was the approach taken by Venter and Şahin (2018).

Another way to determine the degree of shrinkage to use is to put a wide-enough prior distribution on $\lambda$ itself. Then the posterior distribution will include estimation of $\lambda$. Our experience has been that the resulting loo from this ends up close to the highest loo for any $\lambda$, and can be even slightly higher, possibly due to having an entire posterior distribution of $\lambda$s. Gao and Meng (2018) do this, and it is done in the mortality example as well.

Putting a prior on $\lambda$ is considered to be the fully Bayesian approach. Estimating $\lambda$ by cross-validation is not a Bayesian step. Bayesians often think of a parameter search to optimize a cross-validation method as a risky approach. A cross-validation measure like loo is an estimate of how the model would fit a new, independent sample from the same population, but such estimates themselves have estimation errors. A search over parameters to optimize the measure thus runs a risk of producing a measure that is overstated. The fully Bayesian approach avoids this problem. As we saw above, the fully Bayesian method is also fully frequentist, as it can be done entirely by random effects with a postulated distribution for $\lambda$.

2f Postulated shrinkage distributions

The shape of the shrinkage distribution can also have an effect on the model fit. The Laplace distribution is more peaked at zero than the normal, and also has heavier tails, but is lighter at some intermediate values. An increasingly popular shrinkage distribution is the Cauchy

\[
1/p(\beta) = \pi(1 + \lambda^2 \beta^2)/\lambda
\]

\[-\log(p(\beta)) = -\log\lambda + \log\pi + \log(1 + \lambda^2 \beta^2)\]

It is even more concentrated near zero and has still heavier tails than the Laplace. For a
fixed λ, the constants in the density drop out, so the conditional mode minimizes:

\[ NLL + \sum \log(1 + \lambda^2 \beta_j^2) \]

This is an alternative to both lasso and ridge regression. Cauchy shrinkage often produces more parsimonious models than the normal or Laplace do. It can have a bit better or bit worse penalized likelihood, but even if slightly worse, the greater parsimony makes it worth considering. It also seems to produce tighter ranges of parameters.

The Cauchy is the t-distribution with \( \nu = 1 \). If \( \nu = 2 \), the conditional mode minimizes

\[ NLL + 1.5 \sum \log(2 + \lambda^2 \beta_j^2) \]

If \( \nu = 3 \), this becomes:

\[ NLL + 2 \sum \log(3 + \lambda^2 \beta_j^2) \]

The normal is the limiting case of t distributions with ever larger degrees of freedom. Degrees of freedom \( \nu \) does not have to be an integer, so a continuous distribution postulated for \( \nu \) could be used to get an indication for how heavy-tailed the shrinkage distribution should be for a given data set. Actually, the Laplace distribution can be approximated by a t distribution. A t with \( \nu = 6 \) matches a Laplace with scale parameter \( \sqrt{3}/2 \) in both variance and kurtosis (and since the odd moments are zero, matches all 5 moments that exist for this t), so is a reasonable approximation. \( E(1/X, X \neq 0) \) is zero for this t but is undefined for the Laplace, which is due to its sharp point at zero.

We test priors for the example in Section 4. The Laplace is assumed while building the model, but afterwards the shrinkage distribution is tested by using a students-t shrinkage distribution with an assumed prior for \( \nu \), and finding the conditional distribution of \( \nu \) given the data. For this case, the conditional mean is close to \( \nu = 2 \). This is heavier-tailed than the Laplace but less so than the Cauchy. The t-2 distribution, with \( \nu = 2 \), is a closed-form distribution:

\[ f(\beta) = (2 + \beta^2)^{-1.5} \]
\[ F(\beta) = [\beta/\sqrt{2 + \beta^2 + 1}]/2 \]

Still the Cauchy is a viable alternative to get to a more parsimonious model.

The versatility of the t for shrinkage also opens up the possibility of using different but correlated shrinkage for the various parameters, as the multi-variate t prior is an option in the software package we use. For \( \nu = 2 \), the covariances are infinite, but a correlation matrix can
still be input. Two adjacent slope changes are probably quite negatively correlated, as making
one high and one low can often be reversed with little effect on the outcome. Recognizing
this correlation in the assumed shrinkage distribution might improve convergence. We leave
this as a possibility for future research.

2g Semiparametric regression
Harezlak, Ruppert, and Wand (2018) provide an example of using semiparametric methods
to create a curve for mean by year of construction for the floor area a fixed monetary unit
could get for apartments in Warsaw in 2007 – 2009. We fit their data by using Bayesian
shrinkage on the slope changes of linear splines (piecewise linear curves). Figure 1 shows the
data and the two resulting curves, which are quite similar.

Harezlak, Ruppert, and Wand (2018) recommend O’Sullivan penalized cubic splines. These
are a form of smoothing splines that meet a roughness penalty controlled by a smoothing
constant $\lambda$. They pick this $\lambda$ using “generalized cross validation” from Craven and Wahba
(1979). These splines are averages of a number of cubic functions and are not closed form,
but there is available software to calculate them. They also discuss how to pick the number
of knots $K$ for tying together the splines. They feel that with the smoothing done anyway,
35 knots spread out by the quantiles of the data will usually suffice. They do have some
functions for testing the number of knots, however.

Actuarial smoothed linear splines can be done on any variables that can be ordered numerically
1, 2, 3, .... This is natural for years, ages, etc. but can make sense for class variables like
vehicle use classes that have some degree of ranking involved. The parameter shrinkage is
applied to the slope changes between two adjacent linear segments. Initially slope changes
are allowed between all adjacent variables, but if the shrinkage reduces a slope change to
zero, the line segment simply continues at that point, and a knot is eliminated.

Thus the shrinkage chooses the knots, which can then determine how many are needed and
put them where the curves most need to change. This is an advantage over the O’Sullivan
splines, which usually use a fixed number like 35 knots spread according to the number of
observations, not changes in the levels. In our experience, 35 knots is usually enough, but
can sometimes lead to over-parameterization. The degree of shrinkage could be selected by
loo cross validation, which seems as good or better than generalized cross validation once you
have a posterior distribution available, but we use the fully Bayesian method of letting the
conditional distribution of the degree of shrinkage given the data determine this. Bayesians
tend to prefer this approach, as discussed at the end of 2e above.

The key to making this work is that you can create a design matrix with slope-change dummy
variables. The initial design matrix would have dummies for every increment of every variable, like age, period, cohort, etc. The slope change at a point is added to the previous slope, and this slope is added to the next slope change, etc., so the dummy for a slope change increments by 1 at each step. The result is that the variable for year $j$ at an observation from year $i$ will have value $(1 + i - j)_+$, where $x_+$ denotes $\max(0, x)$. With this design matrix and a shrinkage prior for the variables, MCMC creates the curves across each variable while fitting the model.

A useful short-cut to the fitting when there are a lot of variables is to use lasso in an initial sorting out process. The R lasso program glmnet has a built-in cross-validation function cv.glmnet(). It uses its own cross-validation method to select a range of $\lambda$ values that are worth considering. We take the low end of this range, which eliminates the fewest variables, as an initial selection point. Usually more variables are worth taking out, but this can happen later. Adjacent dummy variables are highly negatively correlated, and one can usually substitute for another with little change in the overall fit. This creates difficulties for MCMC, but lasso can quickly sort out the variables with least overall contribution to the model, and then MCMC can readily deal with those that remain. We did this in the Warsaw example and also in the mortality example below. Even if by chance a slope change is left out that would add value to the final model, the impact is likely to be minimal.
An advantage of this method is that the fully Bayesian approach chooses the knots and the smoothing constant as parts of the posterior distributions, and as noted above, fully Bayesian has less risk of producing a model with an exaggerated fit measure than cross validation. Also being able to do this with just a design matrix and MCMC fitting software allows flexibility in model construction. More complex bilinear models like Lee-Carter and Hunt-Blake can be readily fit, as we shall see below. Further, it is straightforward to use any desired parameter distribution, like negative binomial in the example.

3 Joint Mortality Modeling by Shrinkage

We are using data from the Human Mortality Database, which organizes tables by year of death and age at death. The year of birth is approximately year of death minus age at death, depending on whether the death is before or after the birthday in the year. For convenience, we consider cohort to be year of death minus age at death. Cohorts are denoted as \( n \), with parameters \( p[n] \), age at death by \( u \), with parameters \( q[u] \), so the year of death is \( n + u \), with parameters \( r[n + u] \). Also \( c \) is a constant term, that is not shrunk in the modeling. The parameters combine in the models to give the log of the mortality rate. Initially the number of deaths is assumed Poisson in the mortality rate times the exposures, which are the number living at the beginning of the year in the \( n, u \) cell. Theoretically, deaths counts are the sum of Bernoulli processes, and so should be binomial, but for small probabilities, the binomial is very close to the Poisson, which is sometimes easier to model. Later we test the Poisson assumption against heavier-tailed distributions.

A basic model is the linear APC model, which does not include the Lee-Carter trend weights. This has become fairly popular with actuaries and it is our starting point, as it is a purely linear model that can be estimated with standard packages, including lasso packages. Its mortality rate \( m \) given by:

\[
m[n, u] = c + p[n] + q[u] + r[n + u]
\]

With exposures \( e[n, u] \), the deaths \( y[n, u] \) are Poisson in \( e[n, u] \cdot exp(m[n, u]) \). The model of Hunt and Blake (2014), adds cohorts to the model of Lee and Carter (1992), which itself allows the time trend \( r[n + u] \) to vary by age, reflecting the fact that medical advances, etc. tend to improve the mortality rates for some ages more than others. Denoting the trend weight for age \( u \) by \( s[u] \), the Hunt-Blake model we use here is:

\[
m[n, u] = c + p[n] + q[u] + s[u]r[n + u]
\]

Actually Hunt and Blake (2014) allow the possibility of different trends with different age
weights either simultaneously or in succession. Venter and Şahin (2018) suggest an informal test for the need for multiple trends, and find that they indeed help in modeling the US male population for ages 30–89. In the example here we fit the model to ages 50 and up, and did this test, which suggested that one trend is sufficient, so the model is presented in that form. In other populations we have noticed that modeling a wide range of ages over a long period can benefit from multiple trends.

The Hunt-Blake and Lee-Carter models are sometimes described as bilinear APC models, as they are linear if either the $s$ or $r$ parameters are taken as fixed constants. The age weights help model the real phenomenon of differential trends by age, but their biggest weakness is that they are fixed over the observation period, and the ages with the most trend can change over time as medical and public health trends change. The possibility of multiple trends in the Hunt-Blake model can reflect such changes.

To set the basic APC model up to use linear-modeling software packages, the entire array of death counts $y[n, u]$ is typically strung out into a single column. When doing this, it is generally helpful to make parallel columns that record which age, period, and cohort a cell comes from. Then a design matrix can be created where all the variables $p[n], q[u], r[n + u]$ have columns parallel to the death counts, and are represented as 0,1 dummy variables that have a 1 only for the observations that variable applies to. With that, a column vector $\beta$ that has a value for the parameter for each variable can be multiplied on the left by the design matrix to give the fitted value of $m[n, u]$ for each cell. The slope-change design matrix described in Section 2g (excerpted in Table 1) is used later for the semiparametric fits.

For the two population case, assume that the dependent variable $y$ column has all the observations for the first population followed by all the observations for the second population. Our approach is to fit a semiparametric model to the larger population and simultaneously fit a semiparametric model for the differences in the second population from the first. Then the smaller population parameters (in log form) are the sum of the two semiparametric models. With shrinkage applied, the difference parameters hopefully are smaller, so the second population would use fewer effective parameters than the first.

To create the overall design matrix for this, we put the first population’s design matrix in the upper left quadrant, whose columns are the shrinkage dummies for the first population, followed by a matrix of all zeros in the upper right quadrant, which has the variables for the second population. The lower left quadrant repeats the upper left, as the second population starts off with all the parameters of the first population. The same matrix is repeated in the lower right quadrant, here for the differences in the second population’s slope changes from
the first population’s. Hopefully many of these will be small or zero. Then one more column was added for the change in the constant term for the second population. This is all zeros for the first population and all ones for the second population. (The software adds in an overall constant term, which is not shrunk, so a variable for it is not needed.)

MCMC is an effective way to search for good parameters, but these variables make it difficult. Consecutive variables are highly correlated – here many are 99.9% correlated. There could be hundreds of variables in APC models for reasonably large datasets. It could take days or weeks to converge to good parameters running MCMC software on all of those correlated parameters. To speed this up, we start with lasso, which is usually very fast, to identify variables that can be eliminated, and then use a reduced variable set for MCMC. Lasso software, like the R package glmnet, includes cross validation routines to help select $\lambda$. This gives a range of $\lambda$ values that is worth considering. We use the lowest $\lambda$ value this suggests, which leaves the largest number of variables, to generate the starting variables for MCMC. Usually this still includes more parameters than are finally used in the model.

When a slope-change variable is taken out, the slope of the fitted piecewise linear curve does not change at that point, which extends the previous line segment. If instead a small positive or negative value were to be estimated, the slope would change very slightly at that point, probably with not much change to the fitted values. When lasso cross validation suggests that the slope change is not useful, that is saying it would not add to the predictive value of the model. It is possible that a few of the omitted points in the end could have improved the model, but probably they would not have made a lot of difference. So even with MCMC, there is no guarantee that all the possible models have been examined and the very best selected. Nonetheless this procedure is a reasonable way to get very good models.

Constraints are needed to get convergence of APC models. First of all, with the constant term included, the first age, period, and cohort parameters are set to zero. This is not enough to get a unique solution, however. With age, period, and cohort parameters all included, the design matrix is singular. Actuaries use various sets of constraints to deal with this. Using slope change variables, with some eliminated, helps with identifiability. If a straight regression is done for a simple age-period model, the slope change design matrix gives exactly the same $q, r$ parameter values that the 0,1 dummy matrix gives, and the APC matrix is still singular. But when enough slope changes are set to zero, the matrix is no longer singular. Our procedure, then, is to use lasso for an AP design matrix, take out the slope change variables that lasso says are not needed, then add in all the cohort variables, and run lasso again. The maximum variable set lasso finds for this is then the starting point for MCMC, without any other constraints on the variables.
The MCMC runs usually estimate a fair number of the remaining parameters as close to zero. We take out those variables as well, but look at how loo changes. As long as loo does not get worse from this, those variables are left out. Usually a fair number of the variables that lasso identified are eliminated in this process. Leaving them in has little effect on the fit or the loo measure, but the data is more manageable without them. We use the best APC model produced by this as the starting point for the bi-linear models, which add age modifiers to the period trends.

4 Swedish-Danish Male Mortality Example

We model deaths from 1970-2016 for ages 50-99 and cohorts 1883-1953 from the Human Mortality Database. Figure 2 graphs the cumulative mortality trends by age ranges for both populations to see if there are any apparent shifts in the ages with the highest trend. The trends are a bit different in the two countries, but they look a lot like what the Lee-Carter model assumes, with the relative trends among the ages reasonably consistent over time.

4a APC model

Each population has 50 ages for 47 years coming from 71 cohorts. Leaving out the first age, period, and cohort of each population, and adding the dummy variable for Denmark, gives 331 variables for the APC design matrix. Table 1 has an excerpt from the design matrix, showing the first few age and cohort variables for Sweden for several observations.

We started with a lasso run, using the R GLM lasso package glmnet, without the cohort variables. The cv.glmnet app does the lasso estimation as well as this package’s default cross validation. One of the outputs is the smallest \( \lambda \) (thus the one with the most non-zero coefficients) that meets their cross-validation test. We ran the package twice. The first time was without the cohort variables, and then with the resulting non-zero variables from the first run plus all the cohort variables. Below is the key part of the code for that.

```r
fit = cv.glmnet(x, y, standardize = FALSE, family = "poisson", offset=offs)
out <- as.matrix(coef(fit, s = fit$lambda.min))
write.csv(out, file="lasso_fit.csv")
```

This resulted in 80 variables: 22 age variables, 16 year variables, and 42 cohort variables remained after lasso. We use the Stan MCMC application, specifically the R version, rstan. This requires specifying an assumed distribution for each parameter, and it simulates the conditional distribution of the parameters given the data. We used the double exponential distribution for all the slope-change variables. This is a fairly common choice, usually called Bayesian lasso, but we revisit it below. The double exponential distribution in Stan has a scale parameter \( s = 1/\lambda > 0 \). An assumed distribution was provided for that, as well as for
| cohort | year | age | deaths | Sa2 | Sa3 | Sa4 | Sc2 | Sc3 | Sc4 |
|--------|------|-----|--------|-----|-----|-----|-----|-----|-----|
| 1920   | 1970 | 50  | 305    | 0   | 0   | 0   | 37  | 36  | 35  |
| 1919   | 1970 | 51  | 306    | 1   | 0   | 0   | 36  | 35  | 34  |
| 1918   | 1970 | 52  | 344    | 2   | 1   | 0   | 35  | 34  | 33  |
| 1917   | 1970 | 53  | 359    | 3   | 2   | 1   | 34  | 33  | 32  |
| 1916   | 1970 | 54  | 384    | 4   | 3   | 2   | 33  | 32  | 31  |
| 1915   | 1970 | 55  | 417    | 5   | 4   | 3   | 32  | 31  | 30  |
| 1914   | 1970 | 56  | 496    | 6   | 5   | 4   | 31  | 30  | 29  |
| 1913   | 1970 | 57  | 570    | 7   | 6   | 5   | 30  | 29  | 28  |
| 1912   | 1970 | 58  | 569    | 8   | 7   | 6   | 29  | 28  | 27  |
| 1911   | 1970 | 59  | 605    | 9   | 8   | 7   | 28  | 27  | 26  |
| 1910   | 1970 | 60  | 732    | 10  | 9   | 8   | 27  | 26  | 25  |
| 1909   | 1970 | 61  | 781    | 11  | 10  | 9   | 26  | 25  | 24  |
| 1908   | 1970 | 62  | 793    | 12  | 11  | 10  | 25  | 24  | 23  |
| 1907   | 1970 | 63  | 854    | 13  | 12  | 11  | 24  | 23  | 22  |
| 1906   | 1970 | 64  | 953    | 14  | 13  | 12  | 23  | 22  | 21  |
| 1905   | 1970 | 65  | 943    | 15  | 14  | 13  | 22  | 21  | 20  |
| 1904   | 1970 | 66  | 1003   | 16  | 15  | 14  | 21  | 20  | 19  |
| 1903   | 1970 | 67  | 1092   | 17  | 16  | 15  | 20  | 19  | 18  |
| 1902   | 1970 | 68  | 1156   | 18  | 17  | 16  | 19  | 18  | 17  |
| 1901   | 1970 | 69  | 1192   | 19  | 18  | 17  | 18  | 17  | 16  |
| 1900   | 1970 | 70  | 1266   | 20  | 19  | 18  | 17  | 16  | 15  |
| 1899   | 1970 | 71  | 1237   | 21  | 20  | 19  | 16  | 15  | 14  |
| 1898   | 1970 | 72  | 1324   | 22  | 21  | 20  | 15  | 14  | 13  |
| 1897   | 1970 | 73  | 1338   | 23  | 22  | 21  | 14  | 13  | 12  |
| 1896   | 1970 | 74  | 1414   | 24  | 23  | 22  | 13  | 12  | 11  |
| 1895   | 1970 | 75  | 1317   | 25  | 24  | 23  | 12  | 11  | 10  |
| 1894   | 1970 | 76  | 1481   | 26  | 25  | 24  | 11  | 10  | 9   |
| 1893   | 1970 | 77  | 1363   | 27  | 26  | 25  | 10  | 9   | 8   |
| 1892   | 1970 | 78  | 1323   | 28  | 27  | 26  | 9   | 8   | 7   |
| 1891   | 1970 | 79  | 1311   | 29  | 28  | 27  | 8   | 7   | 6   |
| 1890   | 1970 | 80  | 1294   | 30  | 29  | 28  | 7   | 6   | 5   |
| 1889   | 1970 | 81  | 1262   | 31  | 30  | 29  | 6   | 5   | 4   |
| 1888   | 1970 | 82  | 1187   | 32  | 31  | 30  | 5   | 4   | 3   |
| 1887   | 1970 | 83  | 1189   | 33  | 32  | 31  | 4   | 3   | 2   |
| 1886   | 1970 | 84  | 1075   | 34  | 33  | 32  | 3   | 2   | 1   |
| 1885   | 1970 | 85  | 1024   | 35  | 34  | 33  | 2   | 1   | 0   |
| 1884   | 1970 | 86  | 842    | 36  | 35  | 34  | 1   | 0   | 0   |
| 1883   | 1970 | 87  | 764    | 37  | 36  | 35  | 0   | 0   | 0   |
| 1921   | 1971 | 50  | 334    | 0   | 0   | 0   | 38  | 37  | 36  |
| 1920   | 1971 | 51  | 352    | 1   | 0   | 0   | 37  | 36  | 35  |
| 1919   | 1971 | 52  | 330    | 2   | 1   | 0   | 36  | 35  | 34  |
| 1918   | 1971 | 53  | 363    | 3   | 2   | 1   | 35  | 34  | 33  |
Figure 2: Cumulative Trends by Age Group
the constant term.

Eliminating the zero-coefficient variables enabled the APC model to be fit with no further constraints. Usually something like forcing the cohort parameters to have zero trend is needed for identifiability. That particular constraint means that the period trends can be interpreted as the trends given that they have all the trend and none is in the cohorts. The method here is simpler and gives a bit better fit, but does not allow for such an easy interpretation.

For positive parameters, we prefer to start with a distribution proportional to $1/x$. This diverges both as $x$ gets small and as it gets large, so it gives balancing strong pulls up and down, whereas a positive uniform distribution diverges upwardly only, and this can bias the parameter upward. As an example where the integral is known, consider a distribution for the mean $\beta$ of a Poisson distribution with probability function proportional to $e^{-\beta}\beta^k$. With a uniform distribution for $\beta$, which is proportional to 1, the conditional distribution of $\beta$ given an observation $k$ is also proportional to $e^{-\beta}\beta^k$, which makes it a gamma in $k + 1$ and 1, with mean $k + 1$. But if the distribution of $\beta$ is assumed proportional to $1/\beta$, the conditional is proportional to $e^{-\beta}\beta^{k-1}$, which makes it a gamma in $k, 1$, with mean $k$. Thus the $1/\beta$ unconditional distribution take the data at face value, whereas the uniform pushes it upwards. Numerical examples with other distributions find similar results.

The $1/\beta$ assumption is easier to implement in Stan by assuming $log(\beta)$ is uniform. Stan just assumes a uniform distribution is proportional to a constant, so can be ignored. If not specified, it takes the range $\pm 1.7977 \cdot 10^{308}$, which is the largest expressible in double-precision format. We usually start with a fairly wide range for uniform distributions, but if after working with the model for a while the conditional distributions tend to end up in some narrower range, we tend to aim for something a bit wider than that range – mostly to save the program from searching among useless values. We ended up with assuming that the constant is uniform on $[-8,-3]$, with $log(s)$ uniform on $[-6,-3]$.

The best purely APC model had 54 parameters, a loo measure of 22,158.5, and NLL of 22,081.7. The sample-bias penalty, 76.8, is the difference between these. This penalty is a bit high for 54 parameters with shrinkage, which veteran Stan modelers tend to find suspicious in terms of predictive ability of the model. We know from the preliminary visual test that there is some variation in trend across the ages, and not reflecting this could be the problem.

4b Hunt-Blake model

The next step was to put in the trend weights by age, which gives the Hunt-Blake model. This is no longer a simple linear model, which complicates the coding, but not substantially. We used the constraint that the highest weight for any age for a population would be 1.0,
with all the weights positive. In that way, the trend at any year is the trend for the age with
the highest weight.

For this we needed a separate slope-change design matrix for trend weights, with 100 rows
and columns – one for each age in each population. When variables were eliminated, the
columns were dropped but not the rows. Then the parameter vector times the design matrix
gives the weights for each age. In the end, six more age, period, and cohort variables were
eliminated, leaving 46 APC variables, and 72 slope-change variables, so 118 variables all
together. The APC model was refit with the trend weights by age included. This produced a
loo penalized NLL of 20,725.5, an NLL of 20,644.5, which are improvements of about 1400
over the APC model. The penalty of 81.0 for the 118 variables is a more typical relationship
for shrinkage models. NLL and penalized NL are in logs, so it is the absolute difference, not
the relative difference, than indicates how much improvement a better model produces. Here
that is substantial, so Hunt-Blake is clearly better for this data.

4c Alternative shrinkage priors

Next we tried an alternative shrinkage distribution, as discussed in Section 3, using the variables from the best model. Instead of the double
exponential priors, we assumed that the slope-change variables were t-distributed. A double
exponential is roughly equivalent to a t with $\nu = 6$, a Cauchy is $\nu = 1$, and a normal is the
limit as $\nu$ gets large. We tried a prior with $\log(\nu)$ uniform on [-1,6]. That gives a range for $\nu$
of about [0.37,403]. The result was a slightly worse loo, at 20,726. The mean of $\nu$ was 2.4,
with a median of 2. This is heavier tailed than the double exponential, with larger parameters
allowed and smaller ones pushed more towards zero. Two more parameters came out $< 0.001$
and can be eliminated.

We have found in other studies that loo does not change greatly with small changes in the
heaviness of the shrinkage distribution tails, and it looked like estimating $\nu$ was using up
degrees of freedom. Next we tried a constant $\nu = 2$ and took out the two small trend-weight
variables. The result was then 116 variables, with a loo measure of 20,723.4, NLL of 20,643.9,
and a penalty of 79.5. Such a drop of 2 in the penalized NLL is usually considered a better
model, and this one is more parsimonious as well.

4d Negative-binomial model

One assumption not checked so far is the Poisson distribution of deaths in each age-period
cell. As an alternative, we tried the negative binomial distribution, which has an additional
parameter $\phi$, which does not affect the mean but which makes the variance $= \mu^2/\phi + \mu$. This
actually gave a still lower NLL of 20,589.1, and loo of 20,648.1, so an improvement of 75.
The sample-bias penalty was much lower, at 59. There were also three fewer trend-weight
Table 2: Summary of Goodness-of-Fit Measures

| Parameters                                      | Loo Penalty | NLL  |
|-------------------------------------------------|-------------|------|
| APC (Poisson with Laplace shrinkage)            | 22,159      | 22,082 |
| Hunt-Blake (Poisson with Laplace shrinkage)     | 20,726      | 20,645 |
| Hunt-Blake (Poisson with $\nu = 2$ shrinkage)  | 20,723      | 20,644 |
| Hunt-Blake (Negative Binomial with $\nu = 2$ shrinkage) | 20,648      | 20,589 |

Table 3: Parameter Count and Sum of Absolute Values for Negative Binomial Model

| Age | Cohort | Trend | Weight | Sweden Count | Denmark Count | Sweden Sum | Denmark Sum |
|-----|--------|-------|--------|--------------|---------------|------------|-------------|
| 10  | 15     | 5     | 34     | 10           | 3             | 0.126      | 0.022       |
| 3   | 9      | 4     | 33     | 3            | 9             | 0.062      | 0.041       |

variables, leaving 67 of those, as well as 46 age, period, and cohort variables as before. Now there is still a constant, the shrinkage parameter, and $\phi$ that are not shrunk, so 116 variables in total. The low penalty means that the loglikelihood of the omitted variables was not much worse than that of the full model, which is part of why the predictive measure loo was so much better.

The Poisson is an approximation to the binomial distribution that would come from a sum of independent Bernoulli processes. Having a more dispersed distribution suggests that there is some degree of correlation in the deaths. That could arise from high deaths in bad weather or disease outbreaks, for instance. However the dispersion is a measure of the departure of the actual deaths from the fitted means, and also includes model error and estimation error. Thus it is not entirely certain that the better fit implies that deaths are correlated.

We just assumed that $\log(\phi)$ was distributed uniform on $\pm 1.7977 \cdot 10^{308}$. The conditional mean of $\phi$ given the data is 2839. The largest mean for any cell was slightly above 2000, and that would get a variance increased by 70%. This could be as little as a 10% increase for small cells. Compared to the Poisson, this makes it less critical to get a very close fit on the largest cells, which can give a better fit on the other cells. Of course, another reason for the better fit could be actual greater dispersion. The parameters graphed in Figures 3 - 6 are for the negative-binomial model as are the actual vs. fitted values in Figures 7 and 8.

4e Fit summaries

Table 2 summarizes the goodness-of-fit measures for all the models. Table 3 shows the number of parameters of each type by country and the sum of their absolute values in the negative-binomial model. As we were hoping, by doing the joint fitting, Denmark got fewer parameters and lower impact as measured by parameter absolute values. Thus not so much
change from the Sweden model was needed. Trend is a bit of an exception, and as we saw, Sweden does not have much trend while Denmark has some. The mortality-by-age parameters are one area in particular where the populations are similar, and Denmark does not need much change from the Sweden values. It is surprising how dominant the age-weight parameters are in both size and count, and also how they add so little to the NLL penalty and so much to the goodness of fit. They are clearly important in this model.

Figure 3 shows the parametric curves fit to trend factors by population. Sweden has less trend in the later years. Denmark does not deviate from the Swedish trend until halfway through the period. Both show long stretches with few slope changes. These are a lot simpler than the empirical trend graph in Figure 1, which could be why the age weights are important. Figure 4 shows the curves for the age weights. They are more complex than the trends, but still fairly smooth graphs. Denmark largely has the same overall pattern as Sweden, so is using its parameters to a fair degree. Figure 5 graphs the fitted age factors on a log scale, along with a rotation of the factors to horizontal, to better show their differences. There are some nuances of slope-change variation that shows up in the rotation.

Figure 6 shows the cohort factors, along with fitted lines to each to show their average slopes. Although the shapes are different, a lot of the slope changes and differences from linear are actually similar between the two populations.

Actual vs. fitted mortality rates are graphed for the cohorts by age in 3-year ranges in Figure 7. The fitted values form smooth curves in this modeling. Denmark’s rates are more volatile,
but the fitted values reasonably represent the trends in each country. For the most part, the rates are lower for later cohorts. The cohort parameters did not do exactly that, but the fitted rates are a combination of age, period, cohort, and trend-weight parameters, which are all relative to each other. Figure 8 graphs this for year of death instead of cohort, and the conclusions are similar.

5 Conclusion

Joint APC-modeling of related populations is a common need in actuarial work, including for mortality models and loss reserving. Semiparametric regression is a fairly new approach for actuaries, and has advantages over MLE in producing lower estimation and prediction errors. We show how to apply it to individual mortality models by shrinking slope changes of linear fits to the parameters, and then to modeling related data sets by shrinking changes in the semiparametric curves between the data sets.

MCMC estimation can be applied to complex modeling problems. We show how it can be used here to produce samples of the conditional distribution of the parameters given the data. This is typically done in a nominally Bayesian setting, where the model starts with postulated distributions of the parameters, called priors, and yields the conditional distribution of parameters given the data, called the posterior. But frequentist random-effects modeling postulates distributions of the effects (which look like parameters to Bayesians) and there seems to be no reason it could not also use MCMC to sample from the conditional distribution of the effects given the data. Furthermore, when Bayesians take this approach,
Figure 5: Age Factors
they often feel free to depart from the traditional view of priors as incoming beliefs and treat them instead more like random effects – that is postulated distributions. Thus the approach we take appears largely consistent with Bayesian and frequentist approaches, but different from the historical practices of each.

In the example, the difference parameters for the second population are smaller and fewer than the parameters in the first model, suggesting that it is retaining features of the model of the larger population. For this data, the Hunt-Blake model fits better than the linear APC model, the negative binomial fits better than the Poisson, and using the t-distribution with $\nu = 2$ works well as an assumed shrinkage distribution. A more parsimonious model, that looks not as good by loo, can be obtained by a Cauchy shrinkage distribution (the t-distribution with $\nu = 1$), which could be a good idea if the model is regarded as an approximation to a more complex process. These shrinkage distributions also provide minimization formulas similar to those of lasso and ridge regression. The t with $\nu = 2$ case, for instance, leads to minimization of $\text{NLL} + 1.5 \sum_j \log(2 + \lambda^2 \beta_j^2)$.

MCMC software can have problems with highly correlated variables like the slope change variables. Ridge regression and lasso were designed for this problem, and can be much
Figure 7: Actual vs. Fitted Age by Cohort
Figure 8: Actual vs. Fitted Age by Year
more efficient in estimating conditional modes. We use lasso, with only minor shrinkage, to eliminate the less necessary slope-change variables, and then use MCMC with the reduced parameter set. Common lasso software requires linear models, so we start off with a lasso Poisson regression for the linear APC model, with a log link, to reduce the variable set going into MCMC.

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