StMoMo: An R Package for Stochastic Mortality Modelling

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Abstract

In this paper we mirror the framework of generalised (non-)linear models to define the family of generalised Age-Period-Cohort stochastic mortality models which encompasses the vast majority of stochastic mortality projection models proposed to date, including the well-known Lee-Carter and Cairns-Blake-Dowd models. We also introduce the **R** package **StMoMo** which exploits the unifying framework of the generalised Age-Period-Cohort family to provide tools for fitting stochastic mortality models, assessing their goodness of fit and performing mortality projections. We illustrate some of the capabilities of the package by performing a comparison of several stochastic mortality models applied to the England and Wales population.

Keywords: Mortality modelling; mortality forecasting; generalised linear models; generalised non-linear models

1. Introduction

During the last two centuries developed countries experienced a persistent increase in life expectancy. For instance, Oeppen and Vaupel (2002) estimate that during the last 160 years the world record in female life expectancy at birth has increased at an approximate steady pace of 3 months per year. This increase in life expectancy, though a sign of social progress, poses a challenge to governments, private pension plans and life insurers because of its impact on pension and health costs. Actuaries and demographers have recognised the problems caused by an ageing population and rising longevity and have thus devoted significant attention to the development of statistical techniques for the modelling and projection of mortality rates.

One of the most influential approaches to the stochastic modelling of mortality rates is the parsimonious mortality model proposed by Lee and Carter (1992). This model uses principal component analysis to decompose the age-time matrix of mortality rates into a bilinear combination of age and period parameters, with the latter being treated as time series to produce mortality projections. The Lee-Carter model has inspired numerous variants

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and extensions. For instance, Lee and Miller (2001), Booth et al. (2002), and Brouhns et al. (2002) have proposed alternative estimation approaches in order to improve the goodness-of-fit and the forecasting properties of the model. In particular, Brouhns et al. (2002) propose a more formal statistical approach to estimating the parameters by embedding the Lee-Carter model into a Poisson regression setting. Other authors have extended the Lee-Carter model by including additional terms, such as multiple bilinear age-period components (Renshaw and Haberman, 2003; Hyndman and Ullah, 2007), or a cohort effect term (Renshaw and Haberman, 2006; Currie, 2006).

The two factor Cairns-Blake-Dowd (CBD) model introduced by Cairns et al. (2006) is one of the most prominent variants of the Lee-Carter model. The CBD model relies on the linearity of the logit of one-year death probabilities at older ages. Specifically, it assumes that for a given year the logit of the one-year death probability is a linear function of age, and treats the intercept and slope parameters across years as stochastic processes. Cairns et al. (2009) consider three extensions to the original CBD model by incorporating combinations of a quadratic age term and a cohort effect term. Plat (2009) has combined features of the CBD and the Lee-Carter models to produce a model that is suitable for full age ranges and captures the cohort effect.

Given the abundance and rapid increase in the number of stochastic mortality models proposed in the literature, there have been some recent attempts to find the commonalities among these models. Hunt and Blake (2014b) review the structure of mortality models and describe an Age-Period-Cohort model structure which encompasses the vast majority of stochastic mortality models. Currie (2014) shows that many mortality models can be expressed in terms of generalised linear models or generalised non-linear models.

In this paper, we build upon the works of Hunt and Blake (2014b) and Currie (2014) to define the family of Generalised Age-Period-Cohort stochastic mortality models by mirroring the terminology of generalised linear models. We also introduce the **R** package **StMoMo**¹ which exploits the unifying framework of the Generalised Age-Period-Cohort family to provide computational tools for implementing many of the stochastic mortality models proposed to date. The **StMoMo** package is available at https://github.com/amvillegas/StMoMo. Version 0.2.0 has been used for this paper.

Several packages for mortality modelling are available in the **R** environment (R Core Team, 2014). The package **demography** (Hyndman, 2014), whose usage is explained in detail in Booth et al. (2014), implements the original Lee-Carter model along with the Lee and Miller (2001), Booth et al. (2002), and Hyndman and Ullah (2007) variants. The **ilc** package (Butt et al., 2014) implements the Renshaw and Haberman (2006) cohort extension of the Lee-Carter model together with the Lee-Carter model under a Poisson regression framework. The **LifeMetrics R** functions implement the original CBD model and the three

¹The acronym StMoMo, pronounced Saint Momo, stands for Stochastic Mortality Modelling. Momo is the king of Carnivals in numerous Latin American festivities (Wikipedia, 2014).

extended CBD models considered in Cairns et al. (2009), along with the Lee-Carter model (using Poisson maximum likelihood), the Age-Period-Cohort model of Currie (2006) and the Renshaw and Haberman (2006) model. This package, which is not on CRAN, is available at http://www.macs.hw.ac.uk/~andrewc/lifemetrics/.

There are however several drawbacks of the existing packages which our package **StMoMo** seeks to overcome. First, the existing packages are based on model-specific fitting algorithms limiting the models available to those already predefined in the packages. By contrast, **StMoMo** allows users to easily expand the number of models available. In addition, whilst **StMoMo** provides forecasting and simulation functions for any model within the Generalised Age-Period-Cohort family, the existing packages only provide such functions for a limited number of models. For instance, the package **ilc** only includes forecasting functions for the Lee-Carter model. Similarly, simulation with the package **LifeMetrics** is limited to the Lee-Carter and the standard CBD models. Finally, **StMoMo** provides functions which are not available in existing packages, such as tools for analysing the goodness-of-fit² and evaluating the impact of parameter uncertainty using bootstrapping techniques.

StMoMo comes with a set of functions for defining an abstract model — specifying for instance the number of period terms, whether coefficients are parametric or not — and for fitting a given model. This is particularly useful when estimating several models on a given dataset or a given model to different datasets. The package also provides preset functions for defining the most common models available in the mortality forecasting literature. In addition, other models preferred by the user can be created in a very simple fashion, see Section 4 where several examples are given. Therefore, the flexibility of the package allows a user to quickly build up a battery of different models, and this is particularly useful when seeking the most appropriate mortality model, comparing different models or assessing model risk. StMoMo is particularly appealing for actuaries managing life and pensions portfolios exposed to longevity risk. The code backing most functions implemented in the package has been extensively used and tested for the development of multi population mortality models for assessing basis risk in longevity risk transactions, see Baxter et al. (2014).

In the this paper we describe the statistical framework underlying **StMoMo** and illustrate its usage. For this purpose, we use as a running example a comparison of several stochastic mortality models fitted to the England and Wales population. This example is in the spirit of the comparison exercises of Cairns et al. (2009, 2011), Haberman and Renshaw (2011) and Lovász (2011), allowing us to show how several of the analysis performed in these papers can easily be replicated using **StMoMo**. The structure of the paper is as follows. In Section 2 we introduce our notation. In Section 3, we mirror the framework of generalised linear models to define the family of Generalised Age-Period-Cohort (GAPC) stochastic mortality models and demonstrate that many of the mortality models discussed in the literature can be framed within this family. In Section 4 we explain how the GAPC

²We note that **ilc** also provides some graphical tools for assessing the goodness of fit of the models implemented in that package.

family of models is implemented in **StMoMo**. In Section 5, we describe the fitting of GAPC mortality models and illustrate how this can be accomplished using **StMoMo**. In Section 6 we consider the evaluation of the goodness-of-fit of GAPC models. In Section 7 we discuss the forecasting and simulation of GAPC models using time series techniques. Section 8 describes the use of bootstrapping techniques to incorporate parameter uncertainty in the estimation and forecasting of GAPC mortality models. Finally, in Section 9 we provide some conclusions and discuss possible extensions of the **StMoMo** package.

2. Notation and Data

Let the random variable D_{xt} denote the number of deaths in a population at age x last birthday during calendar year t. Also let d_{xt} denote the observed number of deaths, E_{xt}^c the central exposed to risk at age x in year t, and E_{xt}^0 the corresponding initial exposed to risk. The one-year mortality rate for an individual aged x last birthday and in calendar year t, denoted q_{xt} , can be estimated as $\hat{q}_{xt} = d_{xt}/E_{xt}^0$. The force of mortality and central death rates are denoted by μ_{xt} and m_{xt} , respectively, with the empirical estimate of the latter being $\hat{m}_{xt} = d_{xt}/E_{xt}^c$. Under the assumption that the force of mortality is constant over each year of age and calendar year, i.e. from age x to age x+1 and t to t+1, then the force of mortality μ_{xt} and the death rate m_{xt} coincide. We assume that this is the case throughout.

In **StMoMo** and throughout this paper we assume that deaths, d_{xt} , and either central exposures, E_{xt}^c , or initial exposures, E_{xt}^0 , are available in a rectangular array format comprising ages (on the rows) $x = x_1, x_2, \ldots, x_k$, and calendar years (on the columns) $t = t_1, t_2, \ldots, t_n$, When only central exposures are available and initial exposures are required (or vice-versa), one can approximate the initial exposures by adding half the matching reported numbers of deaths to the central exposures, i.e, $E_{xt}^0 \approx E_{xt}^c + \frac{1}{2}d_{xt}$. When the context is clear, we may write E_{xt} to refer to E_{xt}^0 or E_{xt}^c .

3. Generalised Age-Period-Cohort stochastic mortality models

Some authors have recently sought to identify the similarities amongst stochastic mortality models. For instance, Hunt and Blake (2014b) describe an Age-Period-Cohort model structure which encompasses the vast majority of stochastic mortality models. In another interesting contribution, Currie (2014) shows that many common mortality models can be expressed in the standard terminology of generalised linear or non-linear models. In this section, we build upon the aforementioned papers to define the family of Generalised Age-Period-Cohort (GAPC) stochastic mortality models.

Akin to generalised linear models (see e.g. McCullagh and Nelder (1989)), a GAPC stochastic mortality model is comprised of four components:

i. The random component: the numbers of deaths D_{xt} follow a Poisson distribution or a

Binomial distribution³, so that

$$D_{xt} \sim \text{Poisson}(E_{xt}^c \mu_{xt})$$

or

$$D_{xt} \sim \text{Binomial}(E_{xt}^0, q_{xt}),$$

with $\mathbb{E}(D_{xt}/E_{xt}^c) = \mu_{xt}$ and $\mathbb{E}(D_{xt}/E_{xt}^0) = q_{xt}$, respectively.

ii. The systematic component: following Hunt and Blake (2014b) the effects of age x, calendar year t and year-of-birth (cohort) c = t - x are captured through a predictor η_{xt} given by:

$$\eta_{xt} = \alpha_x + \sum_{i=1}^{N} \beta_x^{(i)} \kappa_t^{(i)} + \beta_x^{(0)} \gamma_{t-x}.$$

Here:

- The term α_x is a static age function capturing the general shape of mortality by age.
- $N \geq 0$ is an integer indicating the number of age-period terms describing the mortality trends, with each time index $\kappa_t^{(i)}$, i = 1, ..., N, contributing in specifying the mortality trend and $\beta_x^{(i)}$ modulating its effect across ages.
- The term γ_{t-x} accounts for the cohort effect with $\beta_x^{(0)}$ modulating its effect across ages.

The age modulating terms $\beta_x^{(i)}$, $i=0,1,\ldots,N$, can be either pre-specified functions of age, i.e. $\beta_x^{(i)} \equiv f^i(x)$, as in CBD type models, or non-parametric terms without any prior structure and which need to be estimated as in the Lee-Carter model. In the GAPC family we assume that the period indexes $\kappa_t^{(i)}$, $i=1,\ldots,N$, and the cohort index γ_{t-x} are stochastic processes rather than smooth functions of time or cohort. This is the key feature that allows the stochastic projection of GAPC models and thus the generation of probabilistic forecasts of future mortality rates.

iii. The link function g associating the random component and the systematic component so that

$$g\left(\mathbb{E}\left(\frac{D_{xt}}{E_{xt}}\right)\right) = \eta_{xt}.$$

Although a number of link functions would be possible it is convenient to use the so-called canonical link and pair the Poisson distribution with the log link function and the Binomial distribution with the logit link function (see e.g. Currie (2014) for a discussion of this in the context of mortality models and McCullagh and Nelder (1989) in the wider context of GLMs).

³More precisely, conditionally on (μ_{xt}) , the numbers of deaths are independent and follow a Poisson distribution. Similarly, in the Binomial case, the numbers of deaths are independent conditionally on (q_{xt}) .

iv. The set of parameter constraints: most stochastic mortality models are only identifiable up to a transformation and thus require parameter constraints to ensure unique parameter estimates. These parameter constraints are applied through a constraint function v which maps an arbitrary vector of parameters

$$\theta := \left(\alpha_x, \beta_x^{(1)}, ..., \beta_x^{(N)}, \kappa_t^{(1)}, ..., \kappa_t^{(N)}, \beta_x^{(0)}, \gamma_{t-x}\right)$$

into a vector of transformed parameters

$$v(\theta) = \tilde{\theta} = \left(\tilde{\alpha}_x, \tilde{\beta}_x^{(1)}, ..., \tilde{\beta}_x^{(N)}, \tilde{\kappa}_t^{(1)}, ..., \tilde{\kappa}_t^{(N)}, \tilde{\beta}_x^{(0)}, \tilde{\gamma}_{t-x}\right)$$

satisfying the model constraints with no effect on the predictor η_{xt} (i.e. θ and $\tilde{\theta}$ result in the same η_{xt}).

Most stochastic mortality models proposed in the literature belong to the GAPC family. This includes the original Lee-Carter model, the extensions of the Lee-Carter proposed in Renshaw and Haberman (2003, 2006), the original CBD model, and the extended CBD models of Cairns et al. (2009). In addition, all the model structures considered in Haberman and Renshaw (2011), Lovász (2011) and van Berkum et al. (2014), as well as the models of Plat (2009), Aro and Pennanen (2011), O'Hare and Li (2012), Börger et al. (2013) and Alai and Sherris (2014), are part of the GAPC class of models.⁴

Next, we describe in detail some of these models highlighting how they can be framed within the GAPC family.

3.1. Lee-Carter model under a Poisson setting

The Lee Carter model as implemented by Brouhns et al. (2002) assumes a Poisson distribution of the deaths using a log link function to target the force of mortality μ_{xt} . The predictor structure proposed by Lee and Carter (1992) assumes that there is a static age function, α_x , a unique non-parametric age-period term (N=1), and no cohort effect. Thus, the predictor is given by:

$$\eta_{xt} = \alpha_x + \beta_x^{(1)} \kappa_t^{(1)} \tag{1}$$

In order to project mortality, the time index $\kappa_t^{(1)}$ is modelled and forecasted using ARIMA processes. Typically, a random walk with drift has been shown to provide a reasonable fit, that is,

$$\kappa_t^{(1)} = \delta + \kappa_{t-1}^{(1)} + \xi_t, \qquad \xi_t \sim N(0, \sigma_\kappa),$$

where δ is the drift parameter and ξ_t is Gaussian white noise process with variance σ_{κ} .

⁴We note however that models which rely on the smoothness of mortality over both age and time, such as the graduation approach of Renshaw et al. (1996) and the P-Spline model of Currie (2006), do not belong to the GAPC family. The P-Spline approach is implemented in the **MortalitySmooth** (Camarda, 2012) **R** package.

The Lee-Carter model is only identifiable up to a transformation, as for arbitrary real constants c_1 and $c_2 \neq 0$ the parameters in Equation (1) can be transformed in the following way

$$\left(\alpha_x, \beta_x^{(1)}, \kappa_t^{(1)}\right) \to \left(\alpha_x + c_1 \beta_x^{(1)}, \frac{1}{c_2} \beta_x^{(1)}, c_2(\kappa_t^{(1)} - c_1)\right),\tag{2}$$

leaving η_{xt} unchanged. To ensure identifiability of the model Lee and Carter (1992) suggest the following set of parameter constraints

$$\sum_{x} \beta_x^{(1)} = 1, \qquad \sum_{t} \kappa_t^{(1)} = 0, \tag{3}$$

which can be imposed by choosing

$$c_1 = \frac{1}{n} \sum_t \kappa_t^{(1)}, \qquad c_2 = \sum_x \beta_x^{(1)}$$
 (4)

in transformation (2).

3.2. Renshaw and Haberman model: Lee-Carter with cohort effects

Renshaw and Haberman (2006) generalise the Lee-Carter model by incorporating a cohort effect to obtain the predictor:

$$\eta_{xt} = \alpha_x + \beta_x^{(1)} \kappa_t^{(1)} + \beta_x^{(0)} \gamma_{t-x} \tag{5}$$

Mortality projections for this model are derived using time series forecast of the estimated $\kappa_t^{(1)}$ and γ_{t-x} , generated using univariate ARIMA processes under the assumption of independence between the period and the cohort effects.

To estimate the model Renshaw and Haberman (2006) assume a Poisson distribution of deaths (random component) and use a log link function targeting the force of mortality μ_{xt} . As with the Lee-Carter model, the predictor η_{xt} is invariant to the transformation:

$$\left(\alpha_{x}, \beta_{x}^{(1)}, \kappa_{t}^{(1)}, \beta_{x}^{(0)}, \gamma_{t-x}\right) \to \left(\alpha_{x} + c_{1}\beta_{x}^{(1)} + c_{2}\beta_{x}^{(1)}, \frac{1}{c_{3}}\beta_{x}^{(1)}, c_{3}(\kappa_{t}^{(1)} - c_{1}), \frac{1}{c_{4}}\beta_{x}^{(0)}, c_{4}(\gamma_{t-x} - c_{2})\right),$$

$$(6)$$

where c_1 , c_2 , $c_3 \neq 0$ and $c_4 \neq 0$ are real constants. Identifiability of the model can be ensured using the following set of parameter constraints:

$$\sum_{x} \beta_x^{(1)} = 1, \qquad \sum_{t} \kappa_t^{(1)} = 0, \qquad \sum_{x} \beta_x^{(0)} = 1, \qquad \sum_{c=t_1 - x_k}^{t_n - x_1} \gamma_c = 0,$$

which can be imposed by setting

$$c_1 = \frac{1}{n} \sum_t \kappa_t^{(1)}, \quad c_2 = \frac{1}{n+k-1} \sum_{c=t_1-x_k}^{t_n-x_1} \gamma_c, \quad c_i = \sum_x \beta_x^{(i)}, \quad i = 3, 4,$$

in transformation (6).

Renshaw and Haberman (2006) also consider several substructures of the predictor (5) obtained by setting to a constant one or both of the age modulating terms. Of particular interest is the substructure obtained by setting $\beta_x^{(0)} = 1$,

$$\eta_{xt} = \alpha_x + \beta_x^{(1)} \kappa_t^{(1)} + \gamma_{t-x}, \tag{7}$$

which has been suggested by Haberman and Renshaw (2011) as a simpler structure that resolves some stability issues of the original model.

3.3. Age-Period-Cohort model

Another commonly used substructure of the Renshaw and Haberman model is the so-called Age-Period-Cohort (APC) model, corresponding to $\beta_x^{(1)} = 1$, $\beta_x^{(0)} = 1$,

$$\eta_{xt} = \alpha_x + \kappa_t^{(1)} + \gamma_{t-x},$$

which has a long-standing tradition in the fields of medicine and demography (see, e.g., Clayton and Schifflers (1987) and Hobcraft et al. (1982)), but has not been widely used in the actuarial literature until it was considered by Currie (2006). The APC model is known to be invariant with respect to the following two transformations:

$$\left(\alpha_x, \kappa_t^{(1)}, \gamma_{t-x}\right) \to \left(\alpha_x + \phi_1 - \phi_2 x, \kappa_t^{(1)} + \phi_2 t, \gamma_{t-x} - \phi_1 - \phi_2 (t-x)\right)$$
 (8)

$$\left(\alpha_x, \kappa_t^{(1)}, \gamma_{t-x}\right) \to \left(\alpha_x + c_1, \kappa_t^{(1)} - c_1, \gamma_{t-x}\right),\tag{9}$$

where c_1 , ϕ_1 , and ϕ_2 are real constants. However, we can ensure identifiability of the model by imposing the *set of constraints*:

$$\sum_{t} \kappa_t^{(1)} = 0, \qquad \sum_{c=t_1 - x_k}^{t_n - x_1} \gamma_c = 0, \qquad \sum_{c=t_1 - x_k}^{t_n - x_1} c \gamma_c = 0,$$

where the last two constraints imply that the cohort effect fluctuates around zero with no discernible linear trend. Following Haberman and Renshaw (2011, Appendix A), the constraints on the cohort effect can be imposed by applying transformation (8) with constants ϕ_1 and ϕ_2 obtained by regressing γ_{t-x} on t-x, so that

$$\gamma_{t-x} = \phi_1 + \phi_2(t-x) + \epsilon_{t-x}, \quad \epsilon_{t-x} \sim N(0, \sigma^2)$$
 i.i.d.

The constraint on the period index can then be imposed by applying transformation (9) with

$$c_1 = \frac{1}{n} \sum_t \kappa_t^{(1)}.$$

3.4. CBD model

Cairns et al. (2006) propose a predictor structure with two age-period terms (N=2) with pre-specified age-modulating parameters $\beta_x^{(1)} = 1$ and $\beta_x^{(2)} = x - \bar{x}$, no static age function and no cohort effect. Thus, the predictor of the CBD model is given by:

$$\eta_{xt} = \kappa_t^{(1)} + (x - \bar{x})\kappa_t^{(2)},$$

where \bar{x} is the average age in the data. Cairns et al. (2006) obtain mortality forecasts by projecting the period effects $\kappa_t^{(1)}$ and $\kappa_t^{(2)}$ using a bivariate random walk with drift.

The CBD model does not have identifiability issues and hence the set of parameter constraints is empty. In order to estimate the parameter of the CBD model we can follow Haberman and Renshaw (2011) and assume a Binomial distribution of deaths using a logit link function targeting the one-year death probabilities q_{xt} .

3.5. M7: Quadratic CBD model with cohort effects

Cairns et al. (2009) extend the original CBD model by adding a cohort effect and a quadratic age effect to obtain the predictor:

$$\eta_{xt} = \kappa_t^{(1)} + (x - \bar{x})\kappa_t^{(2)} + \kappa_t^{(3)} \left((x - \bar{x})^2 - \hat{\sigma}_x^2 \right) + \gamma_{t-x},\tag{10}$$

where $\hat{\sigma}_x^2$ is the average value of $(x - \bar{x})^2$. This model, usually referred to as model M7, is not identifiable as the transformation

$$\left(\kappa_t^{(1)}, \kappa_t^{(3)}, \kappa_t^{(3)}, \gamma_{t-x}\right) \to \left(\kappa_t^{(1)} + \phi_1 + \phi_2(t - \bar{x}) + \phi_3\left((t - \bar{x})^2 + \hat{\sigma}^2\right), \kappa_t^{(2)} - \phi_2 - 2\phi_3(t - \bar{x}), \kappa_t^{(3)} + \phi_3, \gamma_{t-x} - \phi_1 - \phi_2(t - x) - \phi_3(t - x)^2\right),$$

$$(11)$$

for real constants ϕ_1 , ϕ_2 and ϕ_3 , leave the predictor unchanged. To identify the model Cairns et al. (2009) impose the set of constraints:

$$\sum_{c=t_1-x_k}^{t_n-x_1} \gamma_c = 0, \qquad \sum_{c=t_1-x_k}^{t_n-x_1} c\gamma_c = 0, \qquad \sum_{c=t_1-x_k}^{t_n-x_1} c^2 \gamma_c = 0,$$

which ensure that the cohort effect fluctuates around zero and has no discernible linear or quadratic trend. Following Haberman and Renshaw (2011, Appendix A), these constraints can be imposed by applying transformation (11) with constants ϕ_1 , ϕ_2 and ϕ_3 obtained by regressing γ_{t-x} on t-x and $(t-x)^2$, so that

$$\gamma_{t-x} = \phi_1 + \phi_2(t-x) + \phi_3(t-x)^2 + \epsilon_{t-x}, \quad \epsilon_{t-x} \sim N(0, \sigma^2)$$
 i.i.d.

Cairns et al. (2009) also consider the simpler predictor structures

$$\eta_{xt} = \kappa_t^{(1)} + (x - \bar{x})\kappa_t^{(2)} + \gamma_{t-x},$$

$$\eta_{xt} = \kappa_t^{(1)} + (x - \bar{x})\kappa_t^{(2)} + (x_c - x)\gamma_{t-x},$$

where x_c is a constant parameter to be estimated. These structures are typically referred to as models M6 and M8, respectively.

3.6. Plat model

Plat (2009) combines the CBD model with some features of the Lee-Carter model to produce a model that is suitable for full age ranges and captures the cohort effect. The proposed predictor structure assumes that there is a static age function, α_x , three age-period terms (N=3) with pre-specified age-modulating parameters $\beta_x^{(1)} = 1$, $\beta_x^{(2)} = \bar{x} - x$, $\beta_x^{(3)} = (\bar{x} - x)^+ = \max(0, \bar{x} - x)$, and a cohort effect with pre-specified age-modulating parameters $\beta_x^{(0)} = 1$. Thus, the predictor is given by:

$$\eta_{xt} = \alpha_x + \kappa_t^{(1)} + (\bar{x} - x)\kappa_t^{(2)} + (\bar{x} - x)^+ \kappa_t^{(3)} + \gamma_{t-x}. \tag{12}$$

Plat (2009) targets the force of mortality μ_{xt} with the log link and estimates the parameters of the model by assuming a Poisson distribution of the deaths. The following parameter transformations leave the predictor in (12) unchanged:

$$\left(\alpha_{x}, \kappa_{t}^{(1)}, \kappa_{t}^{(2)}, \kappa_{t}^{(3)}, \tilde{\gamma}_{t-x}\right) \to \left(\alpha_{x} + \phi_{1} - \phi_{2}x + \phi_{3}x^{2}, \kappa_{t}^{(1)} + \phi_{2}t + \phi_{3}(t^{2} - 2\bar{x}t), \right.$$

$$\left.\kappa_{t}^{(2)} + 2\phi_{3}t, \kappa_{t}^{(3)}, \gamma_{t-x} - \phi_{1} - \phi_{2}(t-x) - \phi_{3}(t-x)^{2}\right) \qquad (13)$$

$$\left(\alpha_{x}, \kappa_{t}^{(1)}, \kappa_{t}^{(2)}, \kappa_{t}^{(3)}, \gamma_{t-x}\right) \to \left(\alpha_{x} + c_{1} + c_{2}(\bar{x} - x) + c_{3}(\bar{x} - x)^{+}, \right.$$

$$\kappa_{t}^{(1)} - c_{1}, \kappa_{t}^{(2)} - c_{2}, \kappa_{t}^{(3)} - c_{3}, \gamma_{t-x}\right), \qquad (14)$$

where c_1 , c_2 , c_3 , ϕ_1 , ϕ_2 , and ϕ_3 are any real constants. The following set of parameter constraints can be imposed to ensure identifiability:

$$\sum_{t} \kappa_{t}^{(1)} = 0, \sum_{t} \kappa_{t}^{(2)} = 0, \sum_{t} \kappa_{t}^{(3)} = 0, \sum_{c=t_{1}-x_{k}}^{t_{n}-x_{1}} \gamma_{c} = 0, \sum_{c=t_{1}-x_{k}}^{t_{n}-x_{1}} c\gamma_{c} = 0, \sum_{c=t_{1}-x_{k}}^{t_{n}-x_{1}} c^{2}\gamma_{c} = 0$$
 (15)

The first three constraints ensure that the period indexes are centred around zero, while the last three constraints ensure that the cohort effect fluctuates around zero and has no linear or quadratic trend. Following Haberman and Renshaw (2011, Appendix A), the constraints on the cohort effect can be imposed by applying transformation (13) with constants ϕ_1 , ϕ_2 , and ϕ_3 obtained by regressing γ_{t-x} on t-x and $(t-x)^2$, so that

$$\gamma_{t-x} = \phi_1 + \phi_2(t-x) + \phi_3(t-x)^2 + \epsilon_{t-x}, \quad \epsilon_{t-x} \sim N(0, \sigma^2) \quad \text{i.i.d.}$$
 (16)

The constraints on the period indexes can then be imposed by applying transformation (14) with

$$c_i = \frac{1}{n} \sum_t \kappa_t^{(i)}, \qquad i = 1, 2, 3.$$
 (17)

In the cases where only older ages are of interest, Plat (2009) suggests to drop the third period term from predictor $(12)^5$:

$$\eta_{xt} = \alpha_x + \kappa_t^{(1)} + (\bar{x} - x)\kappa_t^{(2)} + \gamma_{t-x}.$$
 (18)

We note that this reduced Plat model has the same identifiability issues as the complete model with the omission of the transformations and constraints involving $\kappa_t^{(3)}$ and c_3 .

⁵Note that the reduced Plat model is essentially the M6 model of Cairns et al. (2009) with an added static age term α_x .

4. GAPC stochastic mortality models with StMoMo

The **StMoMo** package provides an **R** implementation of the GAPC family of stochastic mortality models using the standard S3 object-oriented system. The latest development version of **StMoMo** can be installed with the code:

```
install.packages("devtools")
devtools::install_github("amvillegas/StMoMo")
```

The package is loaded within \mathbf{R} as follows:

```
library(StMoMo)
```

In package **StMoMo**, GAPC stochastic mortality models are constructed using the **StMoMo** function. This function takes as input information on the *link function* (and the associated distributional assumption), the *predictor structure*, and the *set of parameter constraints* to create an object of the type "StMoMo" representing a GAPC mortality model:

- The argument link defines the link function and the random component associated with the mortality model. Setting link = "log" assumes that deaths follow a Poisson distribution and uses a log link targeting the force of mortality μ_{xt} , while setting link = "logit" assumes that deaths follow a Binomial distribution and uses a logit link targeting one-year mortality rates q_{xt} .
- The predictor of the model is defined via arguments staticAgeFun, periodAgeFun and cohortAgeFun. Argument staticAgeFun is a logical variable indicating whether the model has a static age function α_x or not. Argument periodAgeFun is a list of length N containing the definitions of the period age-modulating parameters $\beta_x^{(i)}$, $i=1,\ldots,N$, with each entry being either "NP" for non-parametric age terms, "1" for $\beta_x^{(i)}=1$, or a predefined parametric function of age⁶. Argument cohortAgeFun defines the cohort age modulating parameter $\beta_x^{(0)}$ and can take values "NP" for non-parametric age terms, "1" for $\beta_x^{(0)}=1$, a predefined parametric function of age, or NULL if the model does not have a cohort effect.
- The set of parameter constraints are defined via the argument constFun which is a user-defined implementation of the constraint function v mapping an arbitrary vector of parameters to a vector of transformed parameters satisfying the model constraints.

We note that due to limitations of the **R** functions used for fitting "StMoMo" objects to data (see Section 5), the current version **StMoMo** does not support models combining parametric and non-parametric age-modulating functions, $\beta_x^{(i)}$, 0 = 1, ..., N. However, such model are not typically considered and the majority of models proposed in the literature are either extension of the Lee-Carter model with all age-modulating terms being non-parametric or extension of the CBD model with all age-modulating terms being parametric.⁷

⁶Note that we can define a model with no age-period terms (N=0) by making periodAgeFun = NULL.

⁷For instance, a model with predictor structure $\eta_{xt} = \alpha_x + (x - \bar{x})\kappa_t^{(1)} + \beta_x^{(2)}\kappa_t^{(1)}$, corresponding

Table 1: Model structures considered in this paper.

Model	Predictor
LC	$\eta_{xt} = \alpha_x + \beta_x^{(1)} \kappa_t^{(1)}$
CBD	$\eta_{xt} = \kappa_t^{(1)} + (x - \bar{x})\kappa_t^{(2)}$
APC	$\eta_{xt} = \alpha_x + \kappa_t^{(1)} + \gamma_{t-x}$
RH	$\eta_{xt} = \alpha_x + \beta_x^{(1)} \kappa_t^{(1)} + \gamma_{t-x}$
M7	$\eta_{xt} = \kappa_t^{(1)} + (x - \bar{x})\kappa_t^{(2)} + ((x - \bar{x})^2 - \hat{\sigma}_x^2)\kappa_t^{(3)} + \gamma_{t-x}$
PLAT	$\eta_{xt} = \alpha_x + \kappa_t^{(1)} + (\bar{x} - x)\kappa_t^{(2)} + \gamma_{t-x}$

In order to illustrate the creation of particular GAPC mortality models and other capabilities of **StMoMo**, in the rest of this paper we will focus on the models summarised in Table 1. From now onwards, LC stands for the Lee-Carter model; CBD for the original Cairns-Blake-Dowd model; APC for the Age-Period-Cohort model of Currie (2006); RH for the cohort extension of the Lee-Carter model defined in equation (7) and proposed by Renshaw and Haberman (2006); M7 for the quadratic CBD model defined in equation (10); and PLAT for the reduced Plat model defined previously in equation (18). For the sake of comparability, in all cases we will assume a Binomial distribution of deaths and use the logit function to link q_{xt} to the predictor structure η_{xt} .

Below, we show how to define each of the models in Table 1 using the package **StMoMo**.

Lee-Carter model. The LC model under a Binomial setting can be defined using the following code:

Recalling Section 3.1, we note that the constraint function constLC is the R implementation of transformation (2) with constants c_1 and c_2 calculated using Equation (4) to impose the constraints defined in equation (3). The **StMoMo** package also contains the function 1c to

to StMoMo(staticAgeFun = TRUE, periodAgeFun = c(f1, "NP")), with f1 <- function(x, ages) x - mean(ages), is not supported.

facilitate the definition of Lee-Carter models. Hence, we could define the LC model using the much simpler predefined command:

```
LC <- lc(link = "logit")
```

CBD model. To define the CBD model we use the following commands:

Here, we note that function f2 defines the second age-modulating parameter $\beta_x^{(2)} = x - \bar{x}$ and that a constFUN argument needs not be provided since the CBD model does not have identifiability issues. Alternatively, we can define the CBD model using the predefined function cbd:

```
CBD <- cbd()
```

APC model, RH model and model M7. These three models could be defined by implementing explicitly the discussions in Sections 3.2, 3.3 and 3.5. However, **StMoMo** includes predefined functions apc, rh, m7 that facilitate the definition of the APC model, the RH model and model M7, respectively⁸. Thus, these models are defined with the code:

```
RH <- rh(link = "logit", cohortAgeFun = "1")
APC <- apc(link = "logit")
M7 <- m7()</pre>
```

PLAT model. Package **StMoMo** does not include a predefined function for the Plat model. Nevertheless, recalling Section 3.6, we can define the reduced Plat model using the code:

⁸The **StMoMo** package also includes functions m6 and m8 for defining models M6 and M8. We also note that the Renshaw and Haberman (2006) cohort extension of the Lee-Carter in Equation (5) can be defined using function rh with argument cohortAgeFun = "NP".

We note that the constraint function constPlat is the R implementation of transformations (13) and (14) omitting the terms involving $\kappa^{(3)}$ and c_3 , and with constants ϕ_1 , ϕ_2 , ϕ_3 obtained via the linear regression defined in (16) and constant c_1 and c_2 as in Equation (17). Function constPlat imposes the constraints in Equation (15).

5. Model fitting

Parameter estimates of GAPC stochastic mortality models can be obtained by maximising the model log-likelihood, which is given by

$$\mathcal{L}(d_{xt}, \hat{d}_{xt}) = \sum_{x} \sum_{t} \omega_{xt} \left\{ d_{xt} \log \hat{d}_{xt} - \hat{d}_{xt} - \log d_{xt}! \right\}$$

in the case of a Poisson distribution of deaths, and by

$$\mathcal{L}(d_{xt}, \hat{d}_{xt}) = \sum_{x} \sum_{t} \omega_{xt} \left\{ d_{xt} \log \left(\frac{\hat{d}_{xt}}{E_{xt}^{0}} \right) + (E_{xt}^{0} - d_{xt}) \log \left(\frac{E_{xt}^{0} - \hat{d}_{xt}}{E_{xt}^{0}} \right) + \begin{pmatrix} E_{xt}^{0} \\ d_{xt} \end{pmatrix} \right\}$$

in the case of a Binomial distribution of deaths. In both cases, ω_{xt} are weights taking the value 0 if a particular (x, t) data cell is omitted or 1 if the cell is included, and

$$\hat{d}_{xt} = E_{xt} g^{-1} \left(\alpha_x + \sum_{i=1}^N \beta_x^{(i)} \kappa_t^{(i)} + \beta_x^{(0)} \gamma_{t-x} \right)$$

is the expected number of deaths predicted by the model, with g^{-1} denoting the inverse of the link function g.

In the mortality literature, maximisation of the log-likelihood is typically performed using the Newton-Raphson iterative procedure tailored for each model (see e.g. Brouhns et al.

(2002), Renshaw and Haberman (2006) and Cairns et al. (2009)). This is in fact the approach implemented in the packages **ilc** and **LifeMetrics**. Nonetheless, as discussed extensively by Currie (2014), many stochastic mortality models are examples of generalised linear models or generalised non-linear model, which facilitates their fitting using standard statistical software⁹. Currie (2014) exemplifies this fact by fitting several stochastic mortality models in **R** using the standard function glm or the function gnm of the package gnm (Turner and Firth, 2012)¹⁰.

StMoMo provides the generic function fit for estimating the parameters of GAPC mortality models. In line with the remarks of Currie (2014), the corresponding S3 method for objects of the class "StMoMo" relies on function gnm of the package gnm to estimate the parameters of a GAPC model¹¹. Internally, this is accomplished by constructing the equivalent gnm formulation of the GAPC mortality model¹². For instance, the gnm formula of the Binomial LC model created before is

```
LC$gnmFormula
## [1] "D/E ~ -1 + offset(o) + factor(x) + Mult(factor(x), factor(t),
inst = 1)"
```

while the gnm formula of the Binomial CBD model defined before is

```
CBD$gnmFormula

## [1] "D/E ~ -1 + offset(o) + factor(t) + B2:factor(t)"
```

We now illustrate the usage of the function fit of the package StMoMo by fitting the six models defined before to England and Wales mortality data. The object EWMaleData included in the package StMoMo contains data on deaths and central exposures for England and Wales males for the period 1961-2011 and for ages 0-100 obtained from the Human Mortality Database (2014). However, in our examples we concentrate on ages 55 to 89 as the CBD model and the M7 model have been particularly designed to fit higher ages. Additionally, since some models include cohort effects and in agreement with the usual practice (see e.g Cairns et al. (2009) and Haberman and Renshaw (2011)), we exclude (by setting $\omega_{xt} = 0$) all cohorts that have fewer than three observations.

Models LC, APC, CBD, M7 and PLAT can be fitted to England and Wales male mortality data for ages 55 to 89 using the code:

⁹Haberman and Renshaw (2011) have also noticed this fact and profit from GLM facilities in standard statistical packages when fitting CBD type models.

¹⁰Debón et al. (2010) also discuss the use of the package **gnm** for fitting Lee-Carter type models.

¹¹The current version of **StMoMo** only includes the S3 method fit.StMoMo for objects of class "StMoMo", but future versions may provide tailored fit methods for particular stochastic mortality models.

¹²We note that when all the $\beta_x^{(i)}$ are parametric functions of age, the model is a GLM and therefore gnm by default resorts to the glm function of R when fitting the parameter of the model.

```
Dxt <- EWMaleData$Dxt</pre>
Ext <- EWMaleData$Ext + 0.5 * EWMaleData$Dxt
ages <- EWMaleData$ages
years <- EWMaleData$years</pre>
ages.fit <- 55:89
wxt <- genWeightMat(ages = ages.fit, years = years, clip = 3)</pre>
LCfit <- fit(LC, Dxt = Dxt, Ext = Ext, ages = ages, years = years,
             ages.fit = ages.fit, wxt = wxt)
APCfit <- fit(APC, Dxt = Dxt, Ext = Ext, ages = ages, years = years,
             ages.fit = ages.fit, wxt = wxt)
CBDfit <- fit(CBD, Dxt = Dxt, Ext = Ext, ages = ages, years = years,
             ages.fit = ages.fit, wxt = wxt)
M7fit <- fit(M7, Dxt = Dxt, Ext = Ext, ages = ages, years = years,
             ages.fit = ages.fit, wxt = wxt)
PLATfit <- fit(PLAT, Dxt = Dxt, Ext = Ext, ages = ages, years = years,
             ages.fit = ages.fit, wxt = wxt)
```

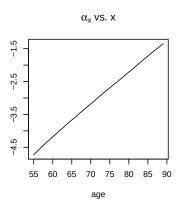
From this code we note the following:

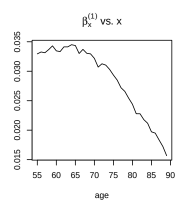
- In order to match the logit-Binomial setting used before in the definition of the mortality models, initial exposures are approximated by transforming the available central exposures.
- The first and last three cohorts years are excluded from the fitting via argument wxt. The appropriate 0-1 weighting matrix, wxt, is constructed using the utility function genWeightMat of package StMoMo.

The fitting of the RH model requires some care as it is well known that fitting cohort extensions of the Lee-Carter models is problematic (Hunt and Villegas, 2014). In particular, Currie (2014) has encountered convergence issues when using package **gnm** to fit the RH model. As a possible way to circumvent these issues, Currie (2014) suggests the use of appropriate starting values when fitting model RH. This can be achieved in function fit via input arguments start.ax, start.bx, start.kt, start.b0x, and start.gc. Using the parameters of the Lee-Carter model as starting values, model RH can be fitted with the code:

The output from the function fit is an object of the class "fitStMoMo" including, among other things, the following information:

- model: the "StMoMo" object defining the underlying GAPC stochastic mortality model;
- ax, bx, kt, b0x, gc: the estimated parameters;





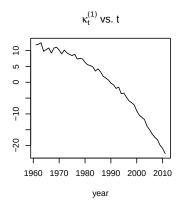


Figure 1: Parameters for the Lee-Carter (LC) model fitted to the England and Wales male population for ages 55-89 and the period 1961-2011.

- loglik: the log-likelihood of the model;
- deviance: the model deviance;
- nobs: the number of observations in the data;
- npar: the effective number of parameters of the model;
- fittingModel: the output of the gnm call used to fit the model.

There are print, plot, fitted, and residuals methods for the "fitStMoMo" class. For instance, Figures 1, 2 and 3 depicting the fitted parameters of the LC model, the CBD model and the APC model, respectively, were produced with the code:

```
plot(LCfit, nCol = 3)
plot(CBDfit, parametricbx = FALSE)
plot(APCfit, parametricbx = FALSE, nCol = 3)
```

6. Goodness-of-fit analysis

The goodness-of-fit of mortality models is typically analysed by inspecting the residuals of the fitted model. Regular patterns in the residuals indicate the inability of the model to describe all the features of the data appropriately. With a Poisson or Binomial random component, it is appropriate to look at the scaled deviance residuals defined as:

$$r_{xt} = \operatorname{sign}(d_{xt} - \hat{d}_{xt}) \sqrt{\frac{\operatorname{dev}(x, t)}{\hat{\phi}}}, \qquad \hat{\phi} = \frac{D(d_{xt}, \hat{d}_{xt})}{K - \nu},$$

where

$$\operatorname{dev}(x,t) = 2d_{xt}\log\left(\frac{d_{xt}}{\hat{d}_{xt}}\right) - (d_{xt} - \hat{d}_{xt})$$

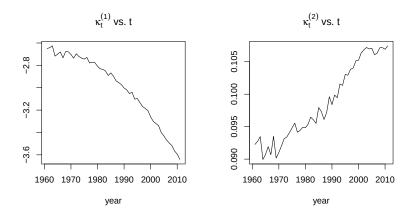


Figure 2: Parameters for the CBD model fitted to the England and Wales male population for ages 55-89 and the period 1961-2011.

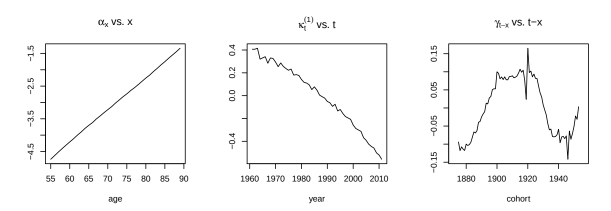


Figure 3: Parameters for the APC model fitted to the England and Wales male population for ages 55-89 and the period 1961-2011.

for a Poisson random component, or

$$dev(x,t) = 2d_{xt} \log \left(\frac{d_{xt}}{\hat{d}_{xt}}\right) + (E_{xt}^0 - d_{xt}) \log \left(\frac{E_{xt}^0 - d_{xt}}{E_{xt}^0 - \hat{d}_{xt}}\right)$$

for a Binomial random component. Further,

$$D(d_{xt}, \hat{d}_{xt}) = \sum_{t} \sum_{t} \omega_{xt} \operatorname{dev}(x, t)$$

is the total deviance of the model, $K = \sum_{x} \sum_{t} \omega_{xt}$ is the number of observations in the data and ν is the effective number of parameters in the model.

In **StMoMo** standardised deviance residuals can be obtained with the generic function residuals applied to a fitted stochastic mortality model of the class "fitStMomo". For example, to obtain the residuals of the LC model and of the CBD model fitted before we use the commands:

```
LCres <- residuals(LCfit)
CBDres <- residuals(CBDfit)</pre>
```

Graphs of these residuals can be produced using the generic function plot. This function supports, via argument type, three types of plots:

- scatter plots of residuals by age, period and cohort such as those extensively used in Haberman and Renshaw (2011);
- black and white sign-plots of the residuals such as those used in Cairns et al. (2009) and Lovász (2011); and
- colour maps (heat-maps) of the residuals.

Figure 4 presents heat-maps of the deviance residuals for the six models fitted to the England and Wales male mortality experience. These charts were produced using function plot with option type = "colourmap". For instance, Figure 4b was obtained with the code:

```
plot(CBDres, type = "colourmap", reslim = c(-3.5, 3.5))
```

From Figure 4 we see that models LC, CBD and APC display strong residual patterns while the residuals of models RH, M7 and PLAT look reasonably random. The APC model shows a strong clustering of residuals due to its inability to allow for varying improvement rates with age. The LC and CBD models, which do not incorporate a cohort effect, show very marked diagonals patterns indicating the inability of these models to capture the well-known cohort effect observed in the England and Wales population (Willets, 2004). The issues with the fit of the LC and CBD models become more evident when looking at scatter plots of the residuals by age, period and cohort. Such plots for the LC model (Figure 5a) and the CBD model (Figure 5b) can be produced using argument type = "scatter" of function plot via the commands:

```
plot(LCres, type = "scatter", reslim = c(-3.5, 3.5))
plot(CBDres, type = "scatter", reslim = c(-3.5, 3.5))
```

The right panels in Figure 5 clearly show that the LC and CBD models are unable to capture the cohort effect. In addition, the left panel in Figure 5b reveals some strong patterns by age, reflecting the the lack of a quadratic age term in the CBD which may be necessary to capture the commonly observed curvature of the mortality rates in a logit scale.

When evaluating the goodness-of-fit of different models, it is generally anticipated that models with more parameters provide a better fit to the data. To rule out the possibility that the better fit observed in a model is the result of over-parametrisation and compare the relative performance of several models, it has become common in the mortality literature to use information criteria which modify the maximum likelihood criterion by penalising models

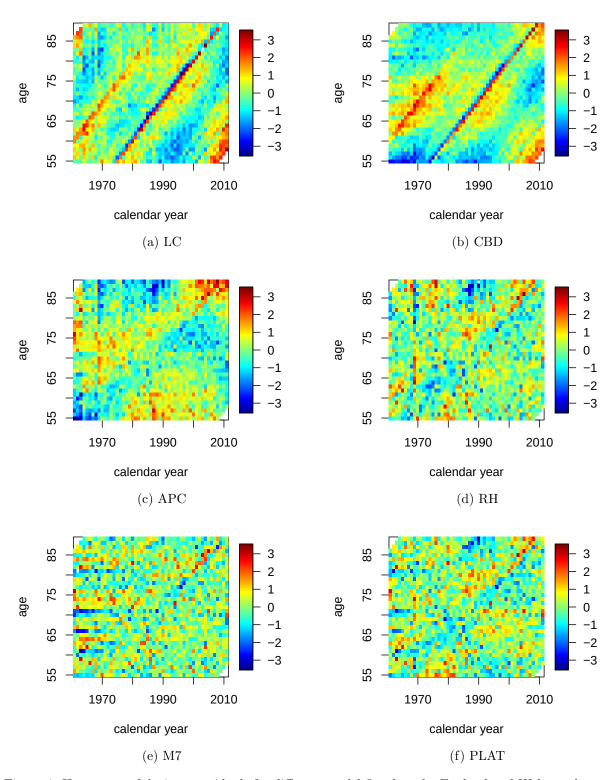


Figure 4: Heat-maps of deviance residuals for different model fitted to the England and Wales males population for ages 55-89 and the period 1961-2011.

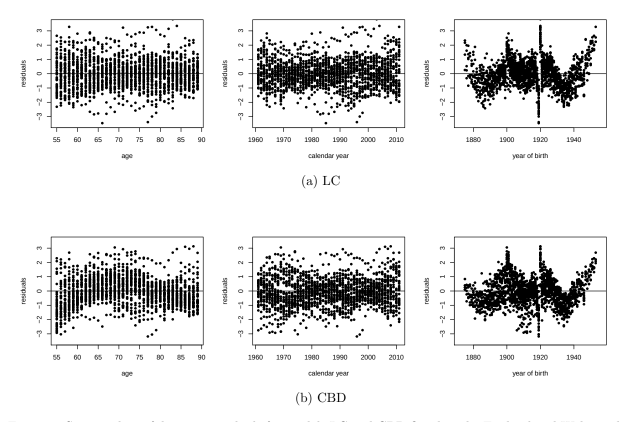


Figure 5: Scatter plots of deviance residuals for models LC and CBD fitted to the England and Wales males population for ages 55-89 and the period 1961-2011.

with more parameters¹³. Two of these criteria are the Akaike Information Criteria (AIC) and the Bayesian Information Criteria (BIC), defined as $AIC = 2\nu - 2\mathcal{L}$ and $BIC = \nu \log K - 2\mathcal{L}$, respectively, with a lower value of AIC and BIC being preferable. In **R** these information criteria can be computed using the generic functions AIC and BIC. For example, we can get the AIC and BIC of the CBD model as follows:

```
## [1] 34697.82

BIC(CBDfit)

## [1] 35256.83
```

Table 2 presents AIC and BIC values for the six models fitted to the England and Wales male data. We note that both criteria lead to the same ranking of models with M7, PLAT, and RH being the best performing models. Overall, these results are consistent with the existing

¹³For examples of such analysis see Cairns et al. (2009, Section 6.1.1), Haberman and Renshaw (2011, Section 3.3), and Lovász (2011, Section 4.1).

-	LC	CBD	APC	RH	M7	PLAT
Number of Parameters	119	102	162	197	229	211
AIC	29866	34698	24469	21779	21406	21624
BIC	30518	35257	25357	22859	22661	22780

Table 2: Number of parameters, AIC and BIC values for different model fitted to the England and Wales males population for ages 55-89 and the period 1961-2011

literature comparing single population models, where the Renshaw-Haberman extension of the Lee-Carter model and the M7 model have been identified as good candidates for modelling mortality in the England and Wales population (Cairns et al., 2009; Haberman and Renshaw, 2011).

7. Forecasting and simulation with stochastic mortality models

In the family of GAPC stochastic mortality models the dynamics of mortality are driven by the period indexes $\kappa_t^{(i)}$, i = 1, ..., N, and the cohort index γ_{t-x} . Therefore, the forecasting and simulation of mortality rates requires the modelling of these indexes using time series techniques.

For the period indexes we follow the standard approach (Cairns et al., 2006, 2011; Haberman and Renshaw, 2011; Lovász, 2011) and use a multivariate random walk with drift. Specifically, we assume that

$$\boldsymbol{\kappa}_{t} = \boldsymbol{\delta} + \boldsymbol{\kappa}_{t-1} + \boldsymbol{\xi}_{t}^{\kappa}, \qquad \boldsymbol{\kappa}_{t} = \begin{pmatrix} \kappa_{t}^{(1)} \\ \vdots \\ \kappa_{t}^{(N)} \end{pmatrix}, \qquad \boldsymbol{\xi}_{t}^{\kappa} \sim N(\mathbf{0}, \Sigma), \tag{19}$$

where $\boldsymbol{\delta}$ is an N-dimension vector of drift parameters and Σ is the $N \times N$ variance-covariance matrix of the multivariate white noise $\boldsymbol{\xi}_t^{\kappa}$.

As pointed out by Currie (2014), the main challenge when forecasting stochastic mortality models is specifying the dynamics of the cohort effect. To have a simple starting point, we follow previous studies (Renshaw and Haberman, 2006; Cairns et al., 2011; Lovász, 2011) and assume that the cohort index, γ_{t-x} , follows a univariate ARIMA process which is independent of the period index, κ_t . In general, we assume that $\gamma_c \equiv \gamma_{t-x}$ follows an ARIMA(p, q, d) with drift, so that

$$\Delta^{d} \gamma_{c} = \delta_{0} + \phi_{1} \Delta^{d} \gamma_{c-1} + \dots + \phi_{p} \Delta^{d} \gamma_{c-p} + \epsilon_{c} + \delta_{1} \epsilon_{c-1} + \dots + \delta_{q} \epsilon_{c-q}, \tag{20}$$

where Δ is the difference operator, δ_0 is the drift parameter, ϕ_1, \ldots, ϕ_p are the autoregressive coefficients with $\phi_p \neq 0, \, \delta_1, \ldots, \delta_q$ are the moving average coefficients with $\delta_q \neq 0$ and ϵ_c is a Gaussian white noise process with variance σ_{ϵ} .

The time series models in (19) and (20) can be used to obtain projected (simulated) values of the period index $\dot{\kappa}_{t_n+s} := \left(\dot{\kappa}_{t_n+s}^{(1)}, \dots, \dot{\kappa}_{t_n+s}^{(N)}\right)'$ and cohort index $\dot{\gamma}_{t_n+s-x_1}$, $s = 1, \dots, h$,

Table 3: ARIMA Models for the cohort effect for models APC, RH, M7 and PLAT.

Mortality Model	Model for γ_{t-x}			
APC	ARIMA(1,1,0) with drift			
RH	ARIMA(1,1,0) with drift			
M7	ARIMA(2,0,0) with non-zero intercept			
PLAT	ARIMA(2,0,0) with non-zero intercept			

respectively, to derive forecasted (simulated) values of the predictor

$$\dot{\eta}_{x,t_n+s} = \alpha_x + \sum_{i=1}^{N} \beta_x^{(i)} \dot{\kappa}_{t_n+s}^{(i)} + \beta_x^{(0)} \dot{\gamma}_{t_n+s-x},$$

which can in turn be used to obtain forecasted (simulated) age-specific central mortality rates, $\dot{\mu}_{x,t_n+s}$ or age-specific one-year mortality rates, \dot{q}_{x,t_n+s} .

In the package **StMoMo** the forecasting of GAPC stochastic mortality models is implemented via the generic method **forecast**. This function estimates and forecasts the multivariate random walk with drift in Equation (19) using the approach described in Haberman and Renshaw (2011, Appendix B)¹⁴ and uses function **Arima** of package **forecast** (Hyndman and Khandakar, 2008) to estimate and forecast the ARIMA process of Equation (20). For instance, if we assume that the cohort indexes of the APC, RH, M7, and PLAT model follow the ARIMA processes specified in Table 3, 50-year ahead (h = 50) central projections of the period indexes, cohort index, and one-year death probabilities for the England and Wales mortality experience can be obtain with the code:

```
LCfor <- forecast(LCfit, h = 50)
CBDfor <- forecast(CBDfit, h = 50)
APCfor <- forecast(APCfit, h = 50, gc.order = c(1, 1, 0))
RHfor <- forecast(RHfit, h = 50, gc.order = c(1, 1, 0))
M7for <- forecast(M7fit, h = 50, gc.order = c(2, 0, 0))
PLATfor <- forecast(PLATfit, h = 50, gc.order = c(2, 0, 0))</pre>
```

The output from the function forecast is an object of the class "forStMoMo" including, among other things, the following information:

• rates: a matrix with the central projection of the mortality rates, $\dot{\mu}_{x,t_n+s}$ or \dot{q}_{x,t_n+s} , $s=1,\ldots,h$;

¹⁴This is implemented in function mrwd of the package **StMoMo**.

- kt.f: a list containing information on the multivariate random walk with drift fitted to the period index κ_t ; and
- gc.f: a list containing information on the ARIMA model fitted to the cohort index γ_{t-x} .

There are print and plot methods for the "forStMoMo" class. For instance, plots of the forecast of the period indexes of models RH, M7 and PLAT (see Figure 6) can be produced using the code:

```
plot(RHfor, only.kt = TRUE)
plot(M7for, only.kt = TRUE, nCol = 3)
plot(PLATfor, only.kt = TRUE)
```

Similarly, plots of the forecast of the cohort indexes of the APC, RH, M7 and PLAT models (see Figure 7) can be obtained with the commands:

```
plot(APCfor, only.gc = TRUE)
plot(RHfor, only.gc = TRUE)
plot(M7for, only.gc = TRUE)
plot(PLATfor, only.gc = TRUE)
```

Package **StMoMo** also provides the function **simulate** for simulating trajectories from GAPC stochastic mortality models. To simulate the period index, κ_t , **StMoMo** implements a multivariate adaptation of Algorithm 2 in Haberman and Renshaw (2009) without provision for parameter error¹⁵, while to simulate the cohort index, γ_{t-x} , function **simulate** uses the equivalent S3 method for objects of class "Arima" provided by the package **forecast**. For example, the code below produces 500 simulated trajectories for the next 50 years of the six stochastic mortality models fitted previously to the England and Wales male mortality experience:

```
set.seed(1234)
nsim <- 500
LCsim <- simulate(LCfit, nsim = nsim, h = 50)
CBDsim <- simulate(CBDfit, nsim = nsim, h = 50)
APCsim <- simulate(APCfit, nsim = nsim, h = 50, gc.order= c(1, 1, 0))
RHsim <- simulate(RHfit, nsim = nsim, h = 50, gc.order= c(1, 1, 0))
M7sim <- simulate(M7fit, nsim = nsim, h = 50, gc.order= c(2, 0, 0))
PLATsim <- simulate(PLATfit, nsim = nsim, h = 50, gc.order= c(2, 0, 0))</pre>
```

¹⁵We note that Algorithm 2 in Haberman and Renshaw (2009) is itself an adaptation of the prediction interval approach of Cairns et al. (2006).

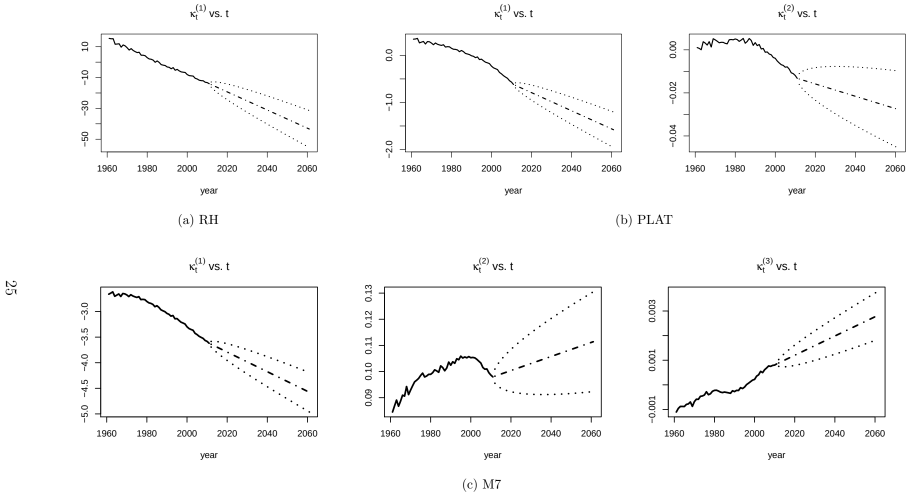


Figure 6: Forecast of the period indexes of the RH, M7 and PLAT models applied to the England and Wales males population for ages 55-89 and the period 1961-2011. Dashed lines represent central forecast and dotted lines represent 95% prediction intervals.

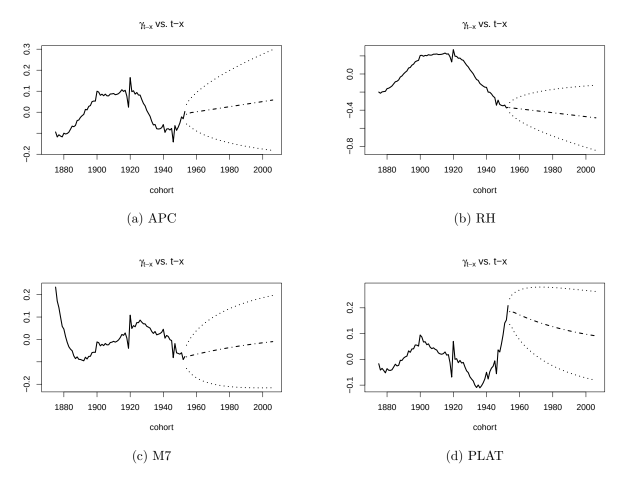


Figure 7: Forecast of the cohort indexes of the APC, RH, M7 and PLAT models applied to the England and Wales males population for ages 55-89 and the period 1961-2011. Dashed lines represent central forecast and dotted lines represent 95% prediction intervals.

The output from the function **simulate** is an object of the class "**simStMoMo**" including, among other things, the following information:

- rates: a three dimensional array with the future simulated mortality rates;
- kt.s: a list containing information on the simulated paths of the period index κ_t ; and
- gc.s: a list containing information on the simulated paths of the cohort index γ_{t-x} .

This output can be used to extract sample trajectories from a model. For instance, Figure 8, which depicts 20 trajectories of the period index, cohort index and one-year death rates at age 65 from model RH, was produced with the code:

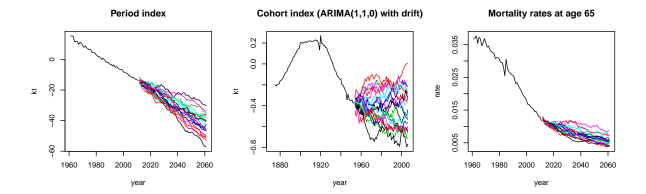


Figure 8: 20 simulated trajectories of the period index $\kappa_t^{(1)}$, cohort index γ_{t-x} and one-year death rates at age 65 q_{xt} for model RH fitted to the England and Wales males population for ages 55-89 and the period 1961-2011.

```
ylim = range(RHfit$kt, RHsim$kt.s$sim[1, , 1:20]),
     type = "l", xlab = "year", ylab = "kt", main = "Period index")
matlines(RHsim$kt.s$years, RHsim$kt.s$sim[1, , 1:20], type = "1", lty = 1)
#Plot cohort index
plot(RHfit$cohorts, RHfit$gc,
     xlim = range(RHfit$cohorts, RHsim$gc.s$cohorts),
     ylim = range(RHfit$gc, RHsim$gc.s$sim[, 1:20], na.rm = TRUE),
     type = "l", xlab = "year", ylab = "kt",
     main = "Cohort index (ARIMA(1,1,0) with drift)")
matlines(RHsim$gc.s$cohorts, RHsim$gc.s$sim[, 1:20], type = "1", lty = 1)
#Plot rates at age 65
qxt <- Dxt / Ext
plot(RHfit$years, qxt["65", ], xlim = range(RHfit$years, RHsim$years),
     ylim = range(qxt["65", ], RHsim$rates["65", , 1:20]), type = "1",
     xlab = "year", ylab = "rate", main = "Mortality rates at age 65")
matlines(RHsim$years, RHsim$rates["65", , 1:20], type = "1", lty = 1)
```

We can also use the output from function simulate to produce fan charts depicting the uncertainty associated with a model forecast. According to Cairns et al. (2011), such plots are central to the analysis of the plausibility of the forecast from a model, and can be used as a criterion when deciding upon what is the most appropriate model amongst a group of possible stochastic mortality models. Figure 9 shows fan charts depicting 50%, 80% and 95% prediction intervals for mortality rates at ages 65, 75 and 85 for each of the six models fitted to the England and Wales experience. Figure 9b, for instance, was produced with the code:

```
library(fanplot)
probs = c(2.5, 10, 25, 50, 75, 90, 97.5)
qxt <- Dxt / Ext</pre>
```

```
#age 65
plot(CBDfit\$years, qxt["65", ], xlim = c(1960, 2061), ylim = c(0.0025, 0.2),
     xlab = "year", ylab = "mortality rate (log scale)",
     pch = 20, log = "y")
fan(t(CBDsim$rates["65", , ]), start = 2012, probs = probs, n.fan = 4,
    fan.col = colorRampPalette(c("black", "white")), ln = NULL)
#age 75
points(CBDfit$years, qxt["75", ], pch = 20)
fan(t(CBDsim$rates["75", , ]), start = 2012, probs = probs, n.fan = 4,
    fan.col = colorRampPalette(c("red", "white")), ln = NULL)
#age 75
points(CBDfit$years, qxt["85", ], pch = 20)
fan(t(CBDsim$rates["85", , ]), start = 2012, probs = probs, n.fan = 4,
    fan.col = colorRampPalette(c("blue", "white")), ln = NULL)
#labels
text(1965, qxt[c("65", "75", "85"), "1990"],
     labels = c("x = 65", "x = 75", "x = 85"))
```

From Figure 9 we note the following:

- Whilst for models CBD, RH, M7 and PLAT the fans at age 85 are wider than at age 65 in accordance with historical evidence (see Cairns et al. (2011, Appendix B)), for models LC and APC the fans at age 85 are narrower than at age 65. This suggest that forecasts from models LC and APC are not plausible for the dataset used in this paper.
- Forecasts for the PLAT model show an implausible increase of mortality rates. This is because the central trend is linked to the estimated cohort effect γ_{t-x} for the PLAT model (see Figure 7d) which shows a steep upward trend between 1935 and 1955.
- The central trend and uncertainty levels produced by the each of the models have noticeably differences. This highlights the importance of recognising model risk as a significant issue when modelling mortality (Cairns et al., 2011).

8. Parameter uncertainty and bootstrapping

When analysing the uncertainty in mortality projections in an actuarial context it is important to consider all sources of risk. However, the prediction intervals (fan charts) obtained in the previous section only account for the uncertainty arising from the error in the forecast of the period and cohort indexes and ignore the uncertainty arising from the estimation of the parameters of the GAPC model.

Due to the analytical intractability of many stochastic mortality models, parameter uncertainty is typically accounted for using the bootstrap procedure. This procedure yields B samples $\alpha_x^b, \beta_x^{(1),b}, \dots, \beta_x^{(N),b}, \kappa_t^{(1),b}, \dots, \kappa_t^{(N),b}, \beta_x^{(0),b}, \gamma_{t-x}^b, \ b=1,\dots,B$, of the parameters

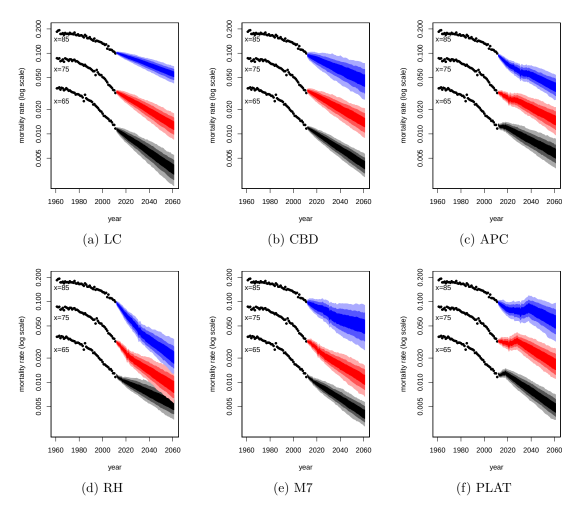


Figure 9: Fan charts for mortality rates q_{xt} at ages x=65 (bottom fan), x=75 (middle fan) and x=85 (top fan) from the six models fitted to the England and Wales males population for ages 55-89 and the period 1961-2011. The dots show historical mortality rates for 1961-2011. Shades in the fan represent prediction intervals at the 50%, 80% and 95% level.

of the GAPC model which can then be used to produce confidence and prediction intervals of demographic and actuarial quantities. To sample the parameters of GAPC models we consider the semiparametric bootstrap proposed by Brouhns et al. (2005) and the residual bootstrap first considered in Koissi et al. (2006):

- Semiparametric bootstrap. Brouhns et al. (2005) propose a semiparametric bootstrap where first B samples of the number of deaths d_{xt}^b , $b = 1, \ldots, B$, are generated by sampling from the Poisson Distribution with mean d_{xt} . Each bootstrapped sample d_{xt}^b , $b = 1, \ldots, B$, is then used to re-estimate the model to obtain B bootstrapped parameter estimates $\alpha_x^b, \beta_x^{(1),b}, \ldots, \beta_x^{(N),b}, \kappa_t^{(1),b}, \ldots, \kappa_t^{(N),b}, \beta_x^{(0),b}, \gamma_{t-x}^b, b = 1, \ldots, B$. Renshaw and Haberman (2008) use the fitted number of death d_{xt} (instead of the observed deaths d_{xt}) to perform the sampling from the Poisson distribution. Wang and Lu (2005) consider a similar semiparametric approach using a Binomial distribution of deaths.
- Residual bootstrap. Another possibility is to bootstrap the residuals of the model

as suggested by Koissi et al. (2006). Under this approach the deviance residuals r_{xt} are resampled with replacement to generate B replications r_{xt}^b , b = 1, ..., B, which are mapped to the corresponding resampled death counts d_{xt}^b , b = 1, ..., B, using the inverse formula. The model is then re-fitted using these resampled number of deaths to produce B sets of estimated parameters α_x^b , $\beta_x^{(1),b}$, ..., $\beta_x^{(N),b}$, $\kappa_t^{(1),b}$, ..., $\kappa_t^{(N),b}$, $\beta_x^{(0),b}$, γ_{t-x}^b , b = 1, ..., B. We refer to Koissi et al. (2006) and Renshaw and Haberman (2008) for details on the inverse formula under a Poisson distribution of deaths and to Debón et al. (2010) for the inverse formula under a Binomial framework. Finally, we note that in implementing the residual bootstrap we follow Renshaw and Haberman (2008) and apply the inverse formula to produce samples of the observed number of deaths rather than samples of the fitted number of deaths as originally done by Koissi et al. (2006).

In what follows we illustrate the assessment of the parameter uncertainty using the package **StMoMo**. In doing so we deviate from the England and Wales example we have used so far and use instead New Zealand mortality data. This new example follows closely the work of Li (2014) who uses New Zealand mortality data to compare several simulation strategies for assessing the risk in mortality projections with a Poisson Lee-Carter model. The main rationale for the change of dataset is that parameter uncertainty is particularly important when analysing the mortality of smaller populations such as smaller countries or pension plans¹⁶. This new example also serves as a means for illustrating the use of the **StMoMo** package with other datasets.

Mortality data for New Zealand can be extracted from the Human Mortality Database (2014) using function hmd.mx of the demography package with the code:

We note that the username and password above are for the Human Mortality Database and should be replaced appropriately. Following Li (2014), we fit a Poisson Lee-Carter model to New Zealand male data for ages 0 to 89 and for the period 1985 to 2008. This can be carried out with the commands:

¹⁶While the population of England and Wales in 2008 was 54.8 million, the population of New Zealand in 2008 was 4.3 million.

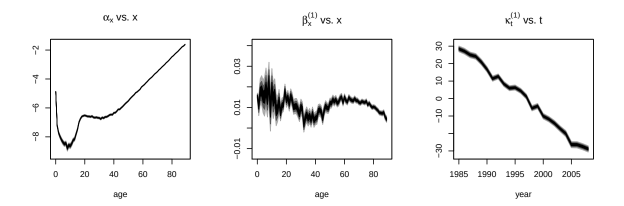


Figure 10: Bootstrapped parameters for the Poisson Lee-Carter model fitted to the New Zealand male population for ages 0-89 and the period 1985-2008. Shades in the fan represent confidence intervals at the 50%, 80% and 95% level.

In the **StMoMo** package the bootstrap of GAPC stochastic mortality models is implemented with the generic function bootstrap. This functions supports both the semiparametric bootstrap and the residual bootstrap. For instance, 5000 semiparametric bootstrap samples of the Lee-Carter model can be obtained with the code¹⁷:

```
LCboot_NZ <- bootstrap(LCfit_NZ, nBoot = 5000, type = "semiparametric")</pre>
```

The output from function bootstrap is an object of the class "bootStMoMo" in which the component bootParameters contains the nBoot replications of the bootstrap parameters. A fan chart depicting the 50%, 80% and 95% confidence intervals of the bootstrapped Lee-Carter model (Figure 10) can be obtained with the command:

plot(LCboot_NZ)

From Figure 10 we note that whilst the parameter uncertainty in the static age function α_x and the period index $\kappa_t^{(1)}$ is modest, the uncertainty in the age-modulating parameters $\beta_x^{(1)}$ is more significant.

Once a stochastic mortality model has been bootstrapped we can simulate it forward to obtain simulated trajectories which account for both the forecast error in the period and cohort indexes and the error in the model fitting. In **StMoMo** we can accomplish this using the function **simulate** applied to an object of class "bootStMoMo". Thus, to obtain 5000 simulated trajectories of the Lee-Carter model for the next 24 years taking into account parameter uncertainty we use the instruction:

 $^{^{17}}$ We note that the bootstrap is a computer intensive procedure. In particular, the 5000 semiparametric bootstrap samples of the Lee-Carter model took about two hours to run in a computer with an Intel Core i5-3320m processor running at 2.60 GHz under Windows 7 Home Premium Edition (64 bits) with 8 GB of RAM.

```
LCsimPU_NZ <- simulate(LCboot_NZ, h = 24)</pre>
```

To highlight the impact of parameter risk on mortality rate projections it is instructive to compare prediction intervals with and without the allowance of parameter uncertainty. 24-year ahead central forecast together with 5000 trajectories of the Lee-Carter model allowing only for forecast error in the random walk with drift and ignoring the model fitting error can be obtained with the code:

```
LCfor_NZ <- forecast(LCfit_NZ, h = 24)
LCsim_NZ <- simulate(LCfit_NZ, nsim = 5000, h = 24)</pre>
```

Figure 11 depicts 95% prediction intervals for mortality rates at age 40, 60 and 80 with and without allowance for parameter uncertainty. This graph was produced with the code:

```
#Observed, fitted and central forecasts
mxt <- LCfit_NZ$Dxt / LCfit_NZ$Ext</pre>
mxtHat <- fitted(LCfit_NZ, type = "rates")</pre>
mxtCentral <- LCfor_NZ$rates</pre>
#95% Prediction intervals without parameter uncertainty
mxtPred2.5 <- apply(LCsim_NZ$rates, c(1, 2), quantile, probs = 0.025)</pre>
mxtPred97.5 <- apply(LCsim_NZ$rates, c(1, 2), quantile, probs = 0.975)</pre>
#95% intervals with parameter uncertainty (in sample, and predictions)
mxtHatPU2.5 <- apply(LCsimPU_NZ$fitted, c(1, 2), quantile, probs = 0.025)</pre>
mxtHatPU97.5 <- apply(LCsimPU_NZ$fitted, c(1, 2), quantile, probs = 0.975)</pre>
mxtPredPU2.5 <- apply(LCsimPU_NZ$rates, c(1, 2), quantile, probs = 0.025)</pre>
mxtPredPU97.5 <- apply(LCsimPU_NZ$rates, c(1, 2), quantile, probs = 0.975)</pre>
#Plot
x \leftarrow c("40", "60", "80")
matplot(LCfit_NZ$years, t(mxt[x, ]),
        xlim = range(LCfit_NZ$years, LCfor_NZ$years),
        ylim = range(mxtHatPU97.5[x, ], mxtPredPU2.5[x, ], mxt[x, ]),
        type = "p", xlab = "years", ylab = "mortality rates (log scale)",
        log = "y", pch = 20, col = "black")
matlines(LCfit_NZ$years, t(mxtHat[x, ]), lty = 1, col = "black")
matlines(LCfit_NZ$years, t(mxtHatPU2.5[x, ]), lty = 5, col = "red")
matlines(LCfit_NZ$years, t(mxtHatPU97.5[x, ]), lty = 5, col = "red")
matlines(LCfor_NZ$years, t(mxtCentral[x, ]), lty = 4, col = "black")
matlines(LCsim_NZ$years, t(mxtPred2.5[x, ]), lty = 3, col = "black")
matlines(LCsim_NZ$years, t(mxtPred97.5[x, ]), lty = 3, col = "black")
matlines(LCsimPU_NZ$years, t(mxtPredPU2.5[x, ]), lty = 5, col = "red")
matlines(LCsimPU_NZ$years, t(mxtPredPU97.5[x, ]), lty = 5, col = "red")
text(1986, mxtHatPU2.5[x, "1995"], labels = c("x=40", "x=60", "x=80"))
```

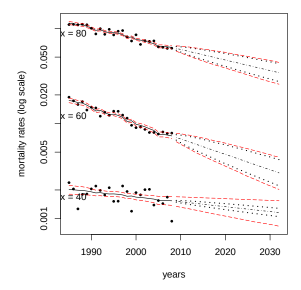


Figure 11: 95% Prediction intervals for mortality rates q_{xt} at ages x=40 (bottom lines), x=60 (middle lines) and x=80 (top lines) for the Poisson Lee-Carter model fitted to the New Zealand male population for ages 0-89 and the period 1985-2008. Dots show historical mortality rates for 1985-2008 and solid black lines show the corresponding fitted rates. Dashed lines represent central forecast, black dotted lines represent 95% prediction intervals excluding parameter uncertainty and dot-dashed red lines depict 95% confidence and prediction intervals including parameter uncertainty.

In Figure 11 we can clearly see that parameter uncertainty has an important impact on the prediction intervals. This is particularly notable at age 40 where in year 2030, for instance, the width of the prediction interval with parameter uncertainty is around 3 times bigger than that without parameter uncertainty. These results are in line with the results obtained by Li (2014) using the same dataset.¹⁸

9. Conclusion

In this paper we have introduced the family of Generalised Age-Period-Cohort stochastic mortality models by paralleling the standard framework of generalised linear models. In addition, we have presented the R package StMoMo which takes advantage of the unifying framework of the GAPC family to provide tools for fitting a diverse number of stochastic mortality models, assessing their goodness of fit and also performing mortality projections. A key feature of the GAPC family and of StMoMo is that they not only encompass models from the Lee-Carter and CBD families, but can also accommodate possible new models. Furthermore, model risk is a prevalent issue when forecasting mortality and we therefore believe that the possibility of easily implementing and comparing a wide range of models makes StMoMo a valuable addition to the toolkit for measuring and managing longevity risk.

 $^{^{18}}$ We note that our prediction intervals without and with parameter uncertainty correspond to methods (I) and (IVa) in Li (2014), respectively.

Our package can be expanded in several directions. The current version of **StMoMo** only allows the use of a log link with a Poisson distribution of deaths or a logit link with a Binomial distribution of deaths. However, we plan to expand the possible combinations of error distribution and link function to include, for instance, Binomial errors with a complementary log-log link as suggested by Currie (2014).

In agreement with the standard in the literature, the **StMoMo** package assumes a multivariate random walk with drift for the dynamics of the period index κ_t . Nevertheless, the package could be expanded to accommodate more general time series models which might be more appropriate for some datasets or some models. For instance, Renshaw and Haberman (2006) use an ARIMA(2,1,0) for $\kappa_t^{(1)}$ when forecasting mortality for England and Wales males with the Lee-Carter model with cohorts, while Plat (2009) uses general univariate ARIMA(p, d, q) for each $\kappa_t^{(i)}$, i = 1, ..., N, fitted using seemingly unrelated regressions to allow for correlations between the different period indexes.

In the GAPC family it is assumed that the age-modulating terms $\beta_x^{(i)}$, i = 0, ..., N, are either non-parametric functions of age which need to be estimated or parametric functions of age with a pre-specified functional form $f^{(i)}(x)$. This latter case could be extended to include the more general case of a pre-specified functional form with a set of parameters that need to be estimated, that is, $\beta_x^{(i)} = f^{(i)}(x; \theta_i)$ with θ_i being some model parameters. This generalisation would allow the implementation of the family of models considered in Hunt and Blake (2014a).

Finally, the increasing attention that multiple population mortality models are receiving in the mortality forecasting literature provides an avenue for a wealth of extensions of our package. In particular, we are considering the development of a two-population version of **StMoMo** that implements the class of relative two-population models considered in Villegas et al. (2015).

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