

Smooth Transition ARMAX Models in Twinkle. (Version 0.9-1

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1 Introduction

The motivation for creating a package for the modelling of smooth transition ARMA models was the observation that many phenomena appear to go through different states which may be explained by some underlying and observable factors.¹ In financial markets, under different states of the business cycle, financial instruments have been observed to exhibit different characteristics with recessions (or the lead up to such) usually marked by increased volatility and lower or negative mean. Being able to model the evolution of the process driving such changes and hence the switch from one state to another must surely make for a better understanding of the underlying dynamics and perhaps lead to a better forecast model.

The precursor to smooth transition models appears to have been Carmichael (1928) who posited the use of the arctangent transformation and even considered the possibility of a double transition well before the plethora of papers which came more than 30 years later. While Quandt (1958) originally discussed a switching regression model, the pioneering contributions to the literature on more general threshold autoregression (TAR) have been Tong and Lim (1980) and Tong and Ghaddar (1981), with a more general class of nonlinear AR models introduced in a series of papers by Billings and Voon (1983), Billings and Voon (1986), and Zhu and Billings (1993). Many extensions to the basic TAR model have been considered in the literature, including the threshold moving average model in Gooijer (1998), threshold ARMA processes in Brockwell et al. (1992), smooth transition based on the Gaussian CDF in Chan and Tong (1986), the more widely adopted logistic (LSTAR) and exponential (ESTAR) models discussed in Teräsvirta (1994), double transitions in ..., nested transition models in Astatkie et al. (1997), multiple states in van Dijk and Franses (1999), and the inclusion of GARCH dynamics in Chan and McAleer (2002, 2003). Thresholds in the variance dynamics have been considered by Zakoian (1994) for GARCH models, and So et al. (2002) for stochastic volatility models. A more general class of hierarchical mixture time series models was proposed by Huerta et al. (2003), while Kalliovirta et al. (2012) proposed a Gaussian mixture AR type model. A type of mixture model is also available in the **twinkle** package and discussed in the next section. For a general review of 'recent' extensions and the state of research in this area see Dijk et al. (2002).

A common theme among the vast majority of econometric research in the area of smooth transition AR models has been the use of the self-exciting model, where the lagged value of the dependent variable (or some simple transformation of the same) is used. The focus on this model has also restricted the representation used in most papers to the one proposed originally by Teräsvirta (1994) which models the state transition rate (γ), or scaling factor, separately from the dynamics. The **twinkle** package departs from this representation and reparameterizes the state dynamics to include a possible linear combination of multivariate variables and the use of autoregressive first order dynamics. The m-states model is also reparameterized to more closely resemble the representation of the multinomial logistic regression model in the way the probabilities are summed and weighted across states. Further extensions include a 2-state AR mixture model partially bridging the gap with the finite mixture models, the inclusion of GARCH dynamics, MA dynamics either inside or outside the states, and a large number of conditional distributions which follow from the rugarch package. Methods for model specification, estimation, filtering, forecasting and simulation are provided with similar interface and access methods as in related packages by the author.

It is important for the interested user to be aware from the start that such models are difficult to estimate, may contain local minima and may be generally hard to solve with confidence. While the package has made efforts to provide for a number of solvers and strategies to estimate these models, confident estimation may prove challenging depending on properties of the dataset used

¹As opposed to unobservable factors which leads to a different class of models, such as the Markov switching models.

and choice of model options. The model is naturally greedy in requiring a substantial amount of data to confidently identify the optimal classification of states given the conditional mean equation. From experience, it is this author's opinion that these types of models may not be as forgiving as linear models when it comes to forecasting, depending on whether the actual forecast state contains the type of nonlinearities under which the model was estimated. Thus, unlike linear models, the misclassification of the forecast state may be more costly.

This paper is organized as follows: Section 2 discusses the representation of the model in the **twinkle** package and how it can be specified. Section 3 discusses forecasting with a special emphasis on the different methods implemented for n-period ahead forecasts, followed by Section ?? on simulation. Finally, Section ?? presents a number of examples using real and simulated data.

While every possible effort has been made to test the model and its methods under different scenarios and squash any bugs, the package is still quite new and further testing is required. As in related packages, a test folder with examples/tests is available in the *inst* folder of the source distribution. The package is available to download from the Bitbucket repository (*alexiosg*) which also contains the development versions of other packages by the author.

General questions on the package should be posted to the R-SIG-FINANCE mailing list, while bugs (with reproducible code and preferably a patch) and suggestions can be reported directly to me.

Finally I would like to acknowledge the valuable help of Eduardo Rossi who collaborated on the new representation and research publication in this area.

2 Smooth Transition ARMA Models Revisited

2.1 Dynamics and Extensions

Consider the standard representation of a 2-state STAR model (adapted from van Dijk and Franses (1999))

$$\begin{aligned}
y_t &= \phi'_1 y_t^{(p)} (F(z_{t-d}; \gamma, \alpha, c)) + \phi'_2 y_t^{(p)} (1 - F(z_{t-d}; \gamma, \alpha, c)) + \varepsilon_t \\
y_t^{(p)} &= \left(1, \tilde{y}_t^{(p)}\right)', \tilde{y}_t^{(p)} = (y_{t-1}, \dots, y_{t-p})' \\
\phi_i &= (\phi_{i0}, \phi_{i1}, \dots, \phi_{ip})' \\
\alpha &= (\alpha_1, \dots, \alpha_k)' \\
\varepsilon_t &\sim ID(0, \sigma) \\
i &= 1, 2(\text{states})
\end{aligned} \tag{1}$$

where ε_t is a white noise zero mean error process with standard deviation σ . The state transition function $F(z_t - 1; \gamma, \alpha, c)$ is a continuous function bounded on the unit interval and usually taken to be the logistic CDF² such that:

$$\begin{aligned}
(\text{Logistic}): F(z_{t-d}; \gamma, \alpha, c) &= (1 + \exp\{-\gamma(\alpha' z_{t-d} - c)\})^{-1}, \gamma > 0 \\
(\text{Exponential}): F(z_{t-d}; \gamma, \alpha, c) &= \left(1 - \exp\left\{-\gamma(\alpha' z_{t-d} - c)^2\right\}\right), \gamma > 0
\end{aligned} \tag{2}$$

where $z_{t-d} = (z_{1t-d}, \dots, z_{jt-d})'$, $j = 1, \dots, k$ is a vector of k observed variables which are assumed to explain the state transition with delay parameter d . These can be a set of explanatory variables or the lagged values of y_t in which case the model is called 'self-exciting'. It is also possible that

²At present only the Logistic STAR model is entertained and it is not likely that the exponential STAR model will be considered at all.

the variable is time in which case the model can be used to identify breaks in the mean as in Lin and Teräsvirta (1994), or a combination of time and other variables giving rise to the time varying STAR (TVSTAR) model discussed in Lundbergh et al. (2003). As correctly noted by van Dijk and Franses (1999), the vector of parameters α needs to be normalized in some way in order to achieve identification (i.e by setting $\alpha_1=1$). The parameter γ is then a type of scaling factor which determines the smoothness (or speed) of the transition, with values at the limits, $[0, \infty]$, representing linear and TAR type transitions respectively. By far the most popular test of STAR nonlinearity is described in Luukkonen et al. (1988) using a Taylor series expansion around equation 2, effectively testing whether $\gamma = 0$ (via a auxilliary regression), which would in turn imply that the α vector is also zero and hence a rejection of STAR type non-linearity. However, there is really little reason for estimating γ seperately in the STAR model since we can allow it to be subsumed by the vector of state parameters (α, c) . In doing so we also gain the additional advantage of extending the type of dynamics to include autoregression as follows:

$$\begin{aligned} F(z_{t-d}; \alpha, c, \beta) &= (1 + \exp\{-\pi_t\})^{-1} \\ \pi_t &= c + \alpha' z_{t-d} + \beta' \pi_t^{(q)} \\ \pi_t^{(q)} &= (\pi_{t-1}, \dots, \pi_{t-q})' \end{aligned} \quad (3)$$

where the unconstrained state dynamics π_t can be initialized by setting

$$\begin{aligned} \pi_0 &= \frac{c + \alpha' \bar{z}}{1 - \beta' \mathbf{1}} \\ \bar{z} &= (E[z_1], \dots, E[z_k])' \end{aligned} \quad (4)$$

with a stationarity constraint of $\left| \sum_{i=1}^q \beta_i \right| < 1$. It should be clear from this that equivalence with the standard representation in Equation 1 is:

$$\begin{aligned} c &= \gamma c \\ \alpha' &= \gamma(1, \alpha_2, \dots, \alpha_j)', j = 1, \dots, k \\ \beta &= 0 \end{aligned} \quad (5)$$

The use of autoregressive dynamics in the state equation follows related work in the area of dynamic binary response models of Kauppi and Saikkonen (2008) and Nyberg (2010). Generally, estimation becomes quite difficult for more than one autoregressive parameter in the state dynamics which is why at present only a lag-1 autoregressive state dynamics model is allowed in the package.

The conditional mean dynamics are not limited to AR terms but may include external regressors (ARX) and moving average (MA) terms in the states giving rise to a full STARMAX model specification:

$$\begin{aligned} y_t &= \left(\phi'_1 y_t^{(p)} + \xi'_1 x_t + \psi'_1 e_t^{(q)} \right) (F(z_{t-d}; \alpha, c, \beta)) \\ &\quad + \left(\phi'_2 y_t^{(p)} + \xi'_2 x_t + \psi'_2 e_t^{(q)} \right) (1 - F(z_{t-d}; \alpha, c, \beta)) + \varepsilon_t \\ \varepsilon_t^{(q)} &= (\varepsilon_{t-1}, \dots, \varepsilon_{t-q})' \\ \psi'_i &= (\psi_{i1}, \dots, \psi_{iq})' \\ x_t &= (x_1, \dots, x_l)' \\ \xi'_1 &= (\xi_{i1}, \dots, \xi_{il})' \end{aligned} \quad (6)$$

It is also possible that the MA term enters outside of the states instead of inside giving rise to a STARX with Linear MA terms (STARXLMA):

$$y_t = \left(\phi'_1 y_t^{(p)} + \xi'_1 x_t \right) (F(z_{t-d}; \alpha, c, \beta)) + \left(\phi'_2 y_t^{(p)} + \xi'_2 x_t \right) (1 - F(z_{t-d}; \alpha, c, \beta)) + \psi' e_t^{(q)} + \varepsilon_t \quad (7)$$

The STARMAX model therefore encompasses a very wide range of sub-models based on the type of restrictions placed in the conditional mean and state dynamics, and choice of switching variables in the latter.

A natural question which arises from the representation is whether it is reasonable to assume that the conditional variance is the same in both states. Consider the STARMAX 2-state model:

$$\begin{aligned} y_t &= \left(\phi'_1 y_t^{(p)} + \xi'_1 x_t + \psi'_1 e_t^{(q)} \right) (F(z_{t-d}; \alpha, c, \beta)) \\ &\quad + \left(\phi'_2 y_t^{(p)} + \xi'_2 x_t + \psi'_2 e_t^{(q)} \right) (1 - F(z_{t-d}; \alpha, c, \beta)) + \varepsilon_t \\ \varepsilon_t &= y_t - (\mu_{1t})p_t - (\mu_{2t})(1 - p_t), d > 0 \end{aligned} \quad (8)$$

Add and subtract $y_t p_t$, and re-arrange:

$$\begin{aligned} \varepsilon_t &= +y_t p_t - (\mu_{1t})p_t + y_t - y_t p_t - (\mu_{2t})(1 - p_t) \\ \varepsilon_t &= +y_t p_t - (\mu_{1t})p_t + y_t (1 - p_t) - (\mu_{2t})(1 - p_t) \\ \varepsilon_t &= (y_t - \mu_{1t})p_t + (y_t - \mu_{2t})(1 - p_t) \\ \varepsilon_t &= (\varepsilon_{1,t})p_t + (\varepsilon_{2,t})(1 - p_t) \\ \varepsilon_{1,t} &\sim N(0, \sigma_1^2) \quad \varepsilon_{2,t} \sim N(0, \sigma_2^2) \\ \varepsilon_t &\sim N(0, \sigma_1^2 p_t + \sigma_2^2 (1 - p_t)) \end{aligned} \quad (9)$$

Thus, the model can naturally be re-formulated as a mixture of Normals³ with mixing probabilities derived from the state dynamics. This provides for a more parsimonious and clear extension than using GARCH dynamics on the mixed state residuals. Alternatively, it can be thought of as the time-invariant version of a STARX-STGARCH model with common transition dynamics. This also provides a partial bridge between finite mixture and time-series autoregressive models.

2.2 Multiple States

van Dijk and Franses (1999) considered an extension of the 2-state STAR model in equation 1 to a 4-state models as follows:

$$\begin{aligned} y_t &= \left[\phi'_1 y_t^{(p)} (1 - F(z_{t-d}; \gamma_1, \alpha, c)) + \phi'_2 y_t^{(p)} (1 - F(z_{t-d}; \gamma_1, \alpha, c)) \right] (1 - F(z_{t-d}; \gamma_2, b, d)) \\ &\quad + \left[\phi'_3 y_t^{(p)} (1 - F(z_{t-d}; \gamma_1, \alpha, c)) + \phi'_4 y_t^{(p)} (1 - F(z_{t-d}; \gamma_1, \alpha, c)) \right] F(z_{t-d}; \gamma_2, b, d) + \varepsilon_t \end{aligned} \quad (10)$$

This effectively models 2 unique states and one interaction state:

$$\begin{aligned} \mu_1 &= \phi'_1 y_t^{(p)} (1 - F(z_{t-d}; \gamma_1, \alpha, c) - F(z_{t-d}; \gamma_2, b, d) + F(z_{t-d}; \gamma_1, \alpha, c) F(z_{t-d}; \gamma_2, b, d)) \\ \mu_2 &= \phi'_2 y_t^{(p)} (1 - F(z_{t-d}; \gamma_1, \alpha, c) - F(z_{t-d}; \gamma_2, b, d) + F(z_{t-d}; \gamma_1, \alpha, c) F(z_{t-d}; \gamma_2, b, d)) \\ \mu_3 &= \phi'_3 y_t^{(p)} (F(z_{t-d}; \gamma_2, b, d) - F(z_{t-d}; \gamma_1, \alpha, c) F(z_{t-d}; \gamma_2, b, d)) \\ \mu_4 &= \phi'_4 y_t^{(p)} (F(z_{t-d}; \gamma_2, b, d) - F(z_{t-d}; \gamma_1, \alpha, c) F(z_{t-d}; \gamma_2, b, d)) \end{aligned} \quad (11)$$

³Or any other location-scale invariant distribution

Alternatively, the implementation followed in this package takes a page out of multinomial regression and models multiple states using the softmax function:

$$y_t = \sum_{i=1}^s \left[\left(\phi'_i y_t^{(p)} + \xi'_i x_t + \psi'_i e_t^{(q)} \right) F_i(z_{t-d}; \alpha_i, c_i, \beta_i) \right] + \varepsilon_t \quad (12)$$

with $s-1$ distinct states modelled:

$$\begin{aligned} F_i(z_{t-d}; \alpha_i, c_i, \beta_i) &= \frac{e^{\pi_{i,t}}}{1 + \sum_{i=1}^{s-1} e^{\pi_{i,t}}} \\ F_s(z_{t-d}; \alpha_i, c_i, \beta_i) &= \frac{1}{1 + \sum_{i=1}^{s-1} e^{\pi_{i,t}}} \end{aligned} \quad (13)$$

where the s states are weighted to sum to unity. This appears, at least to this author, to be a more natural representation for a multi-state setup. In the **twinkle** package, upto 4 states may be modelled⁴

2.3 Estimation

Estimation of the STARMAX models is done by maximizing the likelihood without imposing any particular inequality restrictions on the state dynamic intercepts or any parameter bound restrictions (except for positivity bounds on the variance).⁵ Since unconstrained optimizers appear to do quite well for hard nonlinear/non-smooth problems, the main solver in the **twinkle** package is the BFGS solver from the **optim** function. It is possible to include parameter bounds in which case a logistic transformation is used with the unconstrained solvers. Additional solvers included are 'nlnmb' (bound constrained), 'solnp' (nonlinear SQP solver with nonlinear constraints), 'cmaes' (bound constrained global solver) and 'deoptim' (bound constrained global solver). However, it is suggested that either a multi-start strategy is followed (by choosing 'msoptim') or an iterative search strategy ('strategy') which cycles between fixing the state parameters to estimate the conditional mean parameters (linear), fixing the conditional mean parameters to estimate the state parameters (nonlinear) and a random start estimation. As with general nonlinear optimization problems, scaling of the variables prior to estimation may help, an ica or pca transformation if they are highly correlated, or hinge basis transformation of the dataset via the **earth** package is another interesting option in the case of relevant feature extraction.

3 Forecasting

Consider a general nonlinear first order autoregressive model:

$$y_t = F(y_{t-1}; \theta) + \varepsilon_t \quad (14)$$

where $F(y_{t-1}; \theta)$ is some nonlinear function mapping y_{t-1} to y_t given the parameter set θ . The optimal h -step ahead point forecast, using a least squares criterion, of y_{t+h} at time t is given by:

$$\hat{y}_{t+h|t} = E[y_{t+h} | \mathfrak{F}_t] \quad (15)$$

⁴For the case of only a single state, it is also possible to pass a set of 'time' weights.

⁵Constraints on the autoregressive and moving average parameters may also be placed to impose some type of stationarity constraint per state.

where \mathfrak{S}_t is the information set upto time t . Given that $E[\varepsilon_{t+1} | \mathfrak{S}_t] = 0$, then the 1-step-ahead optimal forecast is:

$$\hat{y}_{t+1|t} = E[y_{t+1} | \mathfrak{S}_t] = F(y_t; \theta) \quad (16)$$

which is the same as when $F(\cdot)$ is linear. However, for horizons greater than 1, this is not the case since $E[F(\cdot)] \neq F(E[\cdot])$, which means that simple recursive relationship found in the linear case do not exist in the nonlinear case. Instead, consider the h-step-ahead point forecast using the following closed form representation:⁶

$$E[y_{t+h} | \mathfrak{S}_t] = \int_{-\infty}^{\infty} E[y_{t+h} | y_{t+h-1}] g(y_{t+h-1} | \mathfrak{S}_t) dy_{t+h-1} \quad (18)$$

where $g(y_{t+h} | \mathfrak{S}_t) = f(y_{t+h} - F(y_{t+h-1}; \theta))$, is the distribution of the shock ε_{t+h} with mean $F(y_{t+h-1})$, though the distribution ε_t is never known with certainty. A number of approaches have been used in the literature to estimate this integral. It is simple to see that the conditional distribution of $g(y_{t+h-1} | \mathfrak{S}_t)$ can be obtained recursively starting at $h=2$ and noting that $g(y_{t+1} | \mathfrak{S}_t) = f(y_{t+1} - F(y_t; \theta))$. To obtain the forecasts, numerical integration can be used (applied recursively) or monte carlo methods. In the former case, the form of the conditional distribution $f(\cdot | \mathfrak{S}_t)$ can be replaced by a kernel estimator, whereas in the latter case one has an option of using an empirical bootstrap, simulating from the conditional distribution $f(\cdot | \mathfrak{S}_t)$ or a kernel estimator. For instance, the 2-step ahead monte carlo forecast is given by:

$$\hat{y}_{t+2|t} = \frac{1}{T} \sum_{i=1}^T F(\hat{y}_{t+1|t} + \varepsilon_i; \theta) \quad (19)$$

However, in the case when a GARCH model is used for the modelling of the conditional variance, then the monte carlo forecast needs to be adjusted as follows:

$$\hat{y}_{t+2|t} = \frac{1}{T} \sum_{i=1}^T F(\hat{y}_{t+1|t} + z_i \hat{\sigma}_{t+2|t}; \theta) \quad (20)$$

where z_i represent draws from either the parametric standardized distribution of the model or the standardized in-sample innovations (or draws from a kernel estimated density of the standardized in-sample innovations), which are then multiplied by the forecast GARCH volatility $\hat{\sigma}_{t+2|t}$ to obtain the forecast residuals ε_i . One benefit of using a monte-carlo or bootstrap approach is that they immediately give rise to the density of each point forecast thus allowing for the creation of interval forecasts.

⁶This is based on the Chapman-Kolmogorov relation:

$$g(y_{t+h} | \mathfrak{S}_t) = \int_{-\infty}^{\infty} g(y_{t+h} | y_{t+h-1}) g(y_{t+h-1} | \mathfrak{S}_t) dy_{t+h-1} \quad (17)$$

which leads to Equation 18 after taking conditional expectations from both sides.

4 Software Implementation

4.1 Specification

The entry point to defining and estimating a STARMAX model in the `twinkle` package is the `starspec` function:

```
>starspec
```

```
function(  
mean.model = list(states=2, include.intercept=c(1,1), arOrder=c(1,1),  
  maOrder=c(0, 0), matype="linear", statevar=c("y","s"), s=NULL,  
  statear=FALSE, ylags=1, xreg=NULL, yfun=NULL, transform="log"),  
variance.model=list(dynamic=FALSE, model="sGARCH", garchOrder=c(1,1),  
  submodel=NULL, vreg=NULL, variance.targeting=FALSE),  
distribution.model="norm", start.pars=list(), fixed.pars=list(),  
fixed.prob=NULL, ...)
```

The **mean.model** defines the equation for the conditional mean dynamics including the state dynamics. Up to 4 **states** are allowed, with the 1-state option having a special implementation in that the **fixed.probs** list is an xts matrix (aligned to the dataset which will be passed to the estimation routine) of weights. By default this is set to a vector of ones in this case but may be any other 'time-weighting' scheme the user wishes. The options for **intercept**, **arOrder** and **maOrder** should be integer vectors of length equal to the number of **states**.

The **matype** denotes whether the moving average terms enter inside the states ('states') or outside ('linear').

The **statevar** indicates whether the model will switch based on its own value ('y') or an exogenous set of regressors ('s'), in which case an xts matrix (aligned to the index of the dataset and appropriately lagged) is passed to **s**. In the case that **statevar** is 'y', then **ylags** is an integer vector of the unique lags to use as a linear combination.

The **yfun** option allows the user to pass a function to transform the value of y^7 prior to being used in the state dynamics equation. While it may appear at first that the same can be achieved by passing a pre-transformed value and using 's' as the **statevar**, consider that simulation and n-ahead forecasts (which depend on simulation methods) on transformed values of 'y' can then be used directly, where it would have been impossible to do so otherwise (because of the path dependency).

The **xreg** is an optional xts matrix of external regressors which needs to be aligned to the index of the dataset (and appropriately lagged). The **statear** indicates whether to include lag-1 autoregression in the state dynamics as discussed previously in equation 3.

The **transform** is currently fixed to use only the logistic transformation, and there are no plans to extend to the exponential at present.

The variance model can be **dynamic**, in which case a choice of 'mixture', 'sGARCH', 'gjrGARCH' and 'eGARCH' are implemented, else by default a static variance model is used. For the GARCH flavors, the rest of the options follow from the `rugarch` package, whilst the 'mixture' model is based on equation 9.

All distributions implemented in `rugarch` are included as options in **distribution.model**, while fixed and starting parameters can be passed directly via **fixed.pars** and **start.pars** respectively, else later on via the **setfixed<-** and **setpars<-** methods on the star specification (note that there is also a **setbounds<-** methods for setting and enforcing parameter bounds).

⁷The function must return the same length as the value it receives without any NAs or NaNs.

Finally, the **fixed.probs** list allows the user to pass an xts matrix of fixed probabilities for each state (aligned to the index of the dataset and with columns equal to **states**). This could for instance be the forecast probabilities from another model (e.g. logistic regression) representing market up and down periods, recessions etc. In this case the state equation is not used and the model is effectively linear and extremely fast to estimate for the conditional mean dynamics.

The returned specification object is of class **STARspec** which may be passed to the estimation routine **starfit**. If the object has been assigned fixed parameters for the complete parameter set, then it may instead be passed to the **starfilter**, **starforecast** or the **starpath** routines.

4.2 Estimation

Once the model has been specified, it may be estimated by maximum likelihood using the **starfit** routine:

```
>starfit

(spec, data, out.sample=0, solver="optim", solver.control=list(),
fit.control=list(stationarity=0, fixed.se=0, rec.init="all"),
cluster=NULL, n=25, ...)
```

The **data** must be an xts object with the same time indices as any data already passed to the **STARspec** object and contain only numeric data without any missing values. The **out.sample** is used to indicate how many data points to optionally leave out in the estimation (from the end of the dataset) for use in out-of-sample forecasting later on when the estimated object is passed to the **starfilter** routine. Perhaps the most important choice to be made is the type of **solver** to use and it's control parameters (**solver.control**). The following solvers and 'strategies' are included:

- **optim**. The preferred choice is the BFGS solver. The choice of solver is controlled by the *method* option in the **solver.control** list.
- **nlminb**. Have had little luck getting the same performance as the BFGS solver.
- **solnp**. Will most likely find a local solution.
- **cmaes**. Even though it is a global solver, it requires careful tweaking of the control parameters (and there are many). This is the parma package version of the solver.
- **deoptim**. Another global solver. May be slow and require tweaking of the control parameters.
- **msoptim**. A multistart version of **optim** with option for using the **cluster** option for parallel evaluation. The number of multi-starts is controlled by the *n.restarts* option in the **solver.control** list.
- **strategy**. A special purpose optimization strategy for STAR problems using the BFGS solver. It cycles between keeping the state variables fixed and estimating the linear variables (conditional mean, variance and any distribution parameters), keeping the linear variables fixed and estimating the state variables, and a random re-start optimization to control for possibly local solutions. The argument **n** in the routine controls the number of times to cycle through this strategy. The **solver.control** list should pass control arguments for the BFGS solver. This is somewhat related to concentrating the sum of squares methodology in terms of the estimation strategy, but does not minimize the sum of squares.

The *strategy* and *msoptim* solver strategies should be the preferred options when estimating STARMA models.

The resulting object of class **STARfit** has a number of methods including an S4 summary (*show*) and a number of S4 extractor methods such as **coef**, **likelihood**, **vcov**, **infocriteria**, **modelmatrix**, **quantile**, **pit**, **fitted**, **residuals** and **sigma**. A new method **states** can be used for extracting the conditional state probabilities, with an extra argument **type** with options for 'prob' (probabilities), 'condm' (conditional mean dynamics per state) and anything else will return the untransformed (raw) state dynamics. The methods are documented in the help page of **STARfit**-class. A default **plot** method is also available which plots the states and fitted values of the model. The package currently does not include any substantial tests for STAR nonlinearity or residual nonlinearity. These may be added in the future.

4.3 Filtering

An object of class **STARspec** with pre-assigned fixed parameter values (for the complete model parameter set) may be passed to the **starfilter** routine with a new or augmented dataset (to that which was used to estimate the original parameters). In this way, new data can be filtered using existing parameters which is equivalent to performing rolling 1-step ahead forecasts (without re-estimation).

```
>starfilter
```

```
(spec, data, out.sample=0, n.old=NULL, rec.init="all", ...)
```

It is probably best to provide an augmented dataset for filtering since the model may depend on the complete history (particularly as regards use of the autoregressive parameter in the state dynamics), in which case the **n.old** option should also be used to denote the size of the original dataset.

4.4 Forecasting

Forecasting can be carried out either from an estimated object of class **STARfit** or a specification with fixed parameters of class **STARspec** (in which case the **data** argument must also be used⁸).

```
>starforecast
```

```
function(fitORspec, data=NULL, n.ahead=1, n.roll=0, out.sample=0,
external.forecasts=list(xregfor=NULL, vregfor=NULL, sfor=NULL,
probfor=NULL), method=c("an.parametric", "an.kernel", "mc.empirical",
"mc.parametric", "mc.kernel"), mc.sims=NULL, ...)
```

As discussed in Section 3, for n-step ($n > 1$) ahead forecasts, there are a number of options available based on recursive quadrature integration ('an') of the integral in Equation 18 else monte carlo ('mc') integration. The *parametric* method uses the density from the estimated object while the *kernel* method fits a kernel density to the residuals. Finally, and only available for the monte carlo method, the *empirical* option samples from the empirical distribution of the residuals. Clearly with a limited history it may be optimal to use either the *parametric* or *kernel* methods. For the monte carlo integration, the **mc.sims** argument denotes the number of samples to use per period.

Some care should be taken when passing **external.forecasts** for the conditional mean regressors (xregfor), the conditional variance regressors (vregfor), the conditional state dynamics

⁸Effectively, the data is filtered with the fixed parameter specification prior to the forecast being carried out

regressors (`sfor`) and the conditional probability (`probfor`) in the case that the state probabilities were passed as fixed in the specification. These xts matrices should be pre-lagged in the same way as the input matrices where in the specification.

The resulting object is of class **STARforecast** with a number of extractor functions documented in the help page of the class and similar to those available for the **STARfit** class. Of particular interest in the case of monte carlo integration is the estimated density of each point forecast which may be extracted from the object.

4.5 Simulation

Simulation can be carried out either directly on a **STARfit** object using the `starsim` routine, else on a **STARspec** object with fixed parameters using the `starp` method.

```
>starsim
```

```
function(fit, n.sim=1000, n.start=0, m.sim=1, presigma=NA, prereturns=NA,
preresiduals=NA, rseed=NA, custom.dist=list(name=NA, distfit=NA),
xregsim=NULL, vregsim=NULL, ssim=NULL, probsim=NULL, ...)
```

The arguments follow similar convention as in related packages. In particular, the **ssim** argument should be a list of matrices for the simulated values of the external regressors (**s**) in the state dynamics, while **probsim** should be a list of matrices of the simulated state probabilities in the case that fixed probabilities were used in the original specification.

Similarly, the `starp` routine has the following arguments:

```
>starp
```

```
function(spec, n.sim=1000, n.start=0, m.sim=1, presigma=NA, prereturns=NA,
preresiduals=NA, rseed=NA, custom.dist=list(name=NA, distfit=NA),
xregsim=NULL, vregsim=NULL, ssim=NULL, probsim=NULL, ...)
```

Where the **prereturns** are now required as depending on the model, so is **presigma** and **preresiduals**. The length of these initialization matrices is determined by the maximum of the conditional state, mean and variance dynamic lags.

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