Particle Filtering with Dynare

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Motivations

- DSGE models are now largely widespread among academic and institutional macro-economists.
- These models are often approximated by a first order Taylor approximation around the steady state.
- The reduced form solution has a linear state-space representation, whose likelihood can be computed with the Kalman filter
- The computational gain comes at the cost that not all questions can be addressed with a linear(ized) model which by nature displays the certainty equivalence property.
- By approximating the model, we also loose some information about the deep parameters to be estimated.
- This paper reviews the econometric approach and their implementation in Dynare for handling DSGE models beyond the (log-)linear approximation.

The reduced form model with a first-order approximation is:

$$z_t = \bar{z}(\Theta) + g_z(\Theta)\hat{z}_{t-1} + g_u(\Theta)u_t$$

which is a (stacked) linear state-space model. Inference on the latent variables and/or parameters is possible with the Kalman filter and MLE.

The reduced form model with a second-order approximation is:

$$\begin{split} z_{t} &= \bar{z}(\Theta) + g_{z}(\Theta)\hat{z}_{t-1} + g_{u}(\Theta)u_{t} \\ &+ 0.5 \, g_{\sigma\sigma}(\Theta) \\ &+ 0.5 \, g_{yy}(\Theta) \, (\hat{z}_{t-1} \otimes \hat{z}_{t-1}) \\ &+ 0.5 \, g_{uu}(\Theta) \, (u_{t} \otimes u_{t}) \\ &+ 0.5 \, g_{uz}(\Theta) \, (\hat{z}_{t-1} \otimes u_{t}) \end{split}$$

Outline of the talk

- Bayesian filtering methods
- Sequential Monte-Carlo methods
- The toy DSGE model
- The filters in Dynare
- Parameters estimation
- **6** Conclusion
- References

The state/space representation

Assume the reduced form of the DSGE model can be cast into the following representation:

$$s_t = f(s_{t-1}, \varepsilon_t; \Theta)$$

$$y_t = g(s_t; \Theta) + e_t$$

In the sequel the sample is denoted $y_{1:T} = \{y_t\}_{t=1}^T$. More generally $x_{1:t} = \{x_i\}_{i=1}^t$ denotes the set of variables x up to time t.

- What can we say about the latent variables? \Rightarrow Characterize the distribution of s_t conditional on the sample up to time t-1 and t (namely $p(s_t|y_{1:t-1})$ and $p(s_t|y_{1:t})$).
- What can we say about the likelihood of an observation? \Rightarrow Characterize the distribution of y_t conditional on the states at time t (s_t) or the sample up to time t-1 (resp. $p(y_t|s_t)$ and $p(y_t|y_{1:t-1})$).

Assumptions

The reduced form DSGE model displays the following properties:

• the first-order Markov relationship on states variables:

$$p(s_t | s_{0:t-1}, y_{1:t-1}) = p(s_t | s_{t-1})$$

unknown but can be easily drawn in by simulating state equations $s_t = f(s_{t-1}, \varepsilon_t; \Theta)$.

• the conditional independence of observations:

$$p(y_t | y_{1:t-1}, s_{0:t}) = p(y_t | s_t)$$

can be (easily) evaluated since measurement errors $e_t (= y_t - g(s_t; \Theta))$ are assumed gaussian.

The theoretical framework

The optimal Bayesian approach

The linear/gaussian case and the Kalman filter

The optimal Bayesian approach

Suppose that the parameters, Θ , and the distribution of the states at time t-1 conditional on the sample up to time t-1, $p(s_{t-1}|y_{1:t-1})$, are known.

Given these informations, we want to characterize how we update our beliefs about the latent variables $(p(s_t|y_{1:t-1}))$ when a new observation y_t becomes available, *i.e.* characterize $p(s_t|y_{1:t})$.

Traditionally, two steps are implemented to obtain this relation:

The prediction step:

$$p(s_t|y_{1:t-1}) = \int p(s_t|s_{t-1}) \ p(s_{t-1}|y_{1:t-1}) ds_{t-1}$$

The updating step:

$$p(s_t|y_{1:t}) = \frac{p(y_t|s_t) p(s_t|y_{1:t-1})}{p(y_t|y_{1:t-1})}$$

where $p(y_t|y_{1:t-1}) = \int p(y_t|s_t) p(s_t|y_{1:t-1}) ds_t$ acts as a constant of integration.

The prediction step

$$p(s_t | y_{1:t-1}) = \int p(s_t, s_{t-1} | y_{1:t-1}) ds_{t-1}$$

$$= \int p(s_t | s_{t-1}, y_{1:t-1}) p(s_{t-1} | y_{1:t-1}) ds_{t-1}$$

$$= \int p(s_t | s_{t-1}) p(s_{t-1} | y_{1:t-1}) ds_{t-1}$$

 $p(s_t | s_{t-1})$ is unknown but can be easily drawn in by simulating state equations $s_t = f(s_{t-1}, \varepsilon_t; \Theta)$.

The updating step

$$p(s_{t}|y_{1:t}) = \frac{p(y_{1:t}|s_{t}) p(s_{t})}{p(y_{1:t})}$$

$$= \frac{p(y_{t}, y_{1:t-1}|s_{t}) p(s_{t})}{p(y_{t}, y_{1:t-1})}$$

$$= \frac{p(y_{t}|y_{1:t-1}, s_{t}) p(y_{1:t-1}|s_{t}) p(s_{t})}{p(y_{t}|y_{1:t-1}) p(y_{1:t-1})}$$

$$= \frac{p(y_{t}|y_{1:t-1}, s_{t}) p(y_{1:t-1}|s_{t}) p(s_{t})}{p(y_{t}|y_{1:t-1}) p(y_{1:t-1})}$$

$$= \frac{p(y_{t}|y_{1:t-1}, s_{t}) p(s_{t}|y_{1:t-1})}{p(y_{t}|y_{1:t-1})}$$

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$$= \frac{p(y_{t}|s_{t}) p(s_{t}|y_{1:t-1})}{p(y_{t}|y_{1:t-1})}$$

 $p(y_t|s_t)$ can be (easily) evaluated since measurement errors are assumed additive and gaussian.

Suppose a linear/gaussian state/space representation:

References

$$s_t = f(s_{t-1}, \varepsilon_t; \Theta) = Ds_{t-1} + E + \varepsilon_t$$

$$y_t = g(s_t; \Theta) + e_t = As_t + B$$

- $p(s_t | y_{1:t-1})$, $p(s_t | y_{1:t})$, $p(y_t | s_t)$ and $p(y_t | y_{1:t-1})$ are all Gaussian.
- Only their conditional expectancy and variance have to be tracked;
- This is what the Kalman filter do analytically:

$$\begin{split} &\mathbb{E}(s_{t}|y_{1:t-1}) = D\mathbb{E}(s_{t-1}|y_{1:t-1}) + E \\ &\mathbb{V}(s_{t}|y_{1:t-1}) = D\mathbb{V}(s_{t-1}|y_{1:t-1})D' + Q \\ &\mathbb{E}(y_{t}|y_{1:t-1}) = A\mathbb{E}(s_{t}|y_{1:t-1}) + B \\ &\mathbb{V}(y_{t}|y_{1:t-1}) = A\mathbb{V}(s_{t}|y_{1:t-1})A' \\ &\mathbb{E}(s_{t}|y_{1:t}) = \mathbb{E}(s_{t}|y_{1:t-1}) + K_{t}[y_{t} - \mathbb{E}(y_{t}|y_{1:t-1})] \\ &\mathbb{V}(s_{t}|y_{1:t}) = (I_{m} - K_{t}A)\mathbb{V}(s_{t}|y_{1:t-1}) \end{split}$$

with $K_t = \mathbb{V}(s_t|y_{1:t-1})A'[\mathbb{V}(y_t|y_{1:t-1})]^{-1}$ the Kalman filter gain.

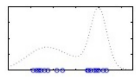
• Otherwise, approximations are required.

Numerical integration (or perfect sampling) Importance sampling Sequential Importance sampling

The likelihood expression

Numerical integration (or perfect sampling)

We assume that the continuous distribution of s_t conditional on $y_{1:t}$ can be approximated by a set of particles $\left\{s_t^{(i)}\right\}_{i=1:N}$ with weights $\left\{w_t^{(i)}\right\}_{i=1:N}$ (summing to one).



Any moments of this distribution can be approximated by a weighted average as follows:

$$\mathbb{E}_{p(s_{t}|y_{1:t})} [h(s_{t})] = \int h(s_{t}) p(s_{t}|y_{1:t}) ds_{t}$$

$$\approx \sum_{i=1}^{N} w_{t}^{(i)} h\left(s_{t}^{(i)}\right) = \frac{1}{N} \sum_{i=1}^{N} h\left(s_{t}^{(i)}\right)$$

Numerical integration (or perfect sampling)
Importance sampling
Sequential Importance sampling
Degeneracy problem and resampling
Summary
The likelihood expression

The idea

Problem: $p(s_t|y_{1:t})$ is unknown.

Solution: **importance sampling**, *i.e.* choose an easy-to-sample *proposal* distribution $q(s_t | y_{1:t})$ (informative on the target distribution)

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$$\mathbb{E}_{p(s_t|y_{1:t})} [h(s_t)] = \int h(s_t) \frac{p(s_t|y_{1:t})}{q(s_t|y_{1:t})} q(s_t|y_{1:t}) ds_t$$

$$= \mathbb{E}_{q(s_t|y_{1:t})} [\tilde{w}_t(s_t)h(s_t)]$$

with normalized weights defined as:

$$\tilde{w}_t(s_t) \equiv \frac{p(s_t|y_{1:t})}{q(s_t|y_{1:t})}$$

These weights can be viewed as importance ratios. Skip details

Numerical integration (or perfect sampling Importance sampling Sequential Importance sampling Degeneracy problem and resampling Summary The likelihood expression

Importance sampling

We can equivalently rewrite these weights, by reversing the conditioning in the numerator with the Bayes theorem, as:

$$\tilde{w}_t(s_t) \equiv \frac{p(y_{1:t}|s_t)p(s_t)}{p(y_{1:t})q(s_t|y_{1:t})}$$

Removing the constant of integration (with respect to s_t), we define the *unormalized* weights:

$$\hat{w}_t(s_t) \equiv \frac{p(y_{1:t}|s_t)p(s_t)}{q(s_t|y_{1:t})}$$

A modification that avoids the calculation of $p(y_{1:t})$:

$$\mathbb{E}_{p(s_t|y_{1:t})}[h(s_t)] = \frac{\mathbb{E}_{q(s_t|y_{1:t})}[\hat{w}_t(s_t)h(s_t)]}{\mathbb{E}_{q(s_t|y_{1:t})}[\hat{w}_t(s_t)]}$$

with unormalized weights defined as:

$$\hat{w}_t(s_t) \equiv \frac{p(y_{1:t}|s_t) p(s_t)}{q(s_t|y_{1:t})} \propto \tilde{w}_t(s_t) \equiv \frac{p(y_{1:t}|s_t) p(s_t)}{q(s_t|y_{1:t}) p(y_{1:t})}$$

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Nonlinear estimation with Dynare

Importance sampling

$$\tilde{w}_t(s_t) \equiv \frac{p(s_t \mid y_{1:t})}{q(s_t \mid y_{1:t})} = \frac{p(y_{1:t} \mid s_t) p(s_t)}{q(s_t \mid y_{1:t}) p(y_{1:t})} \propto \frac{p(y_{1:t} \mid s_t) p(s_t)}{q(s_t \mid y_{1:t})} \equiv \hat{w}_t(s_t)$$

A modification that avoids the calculation of $p(y_{1:t})$:

$$\begin{aligned} p(y_{1:t}) &= \int \frac{p(y_{1:t} \mid s_t) \, p(s_t)}{q(s_t \mid y_{1:t})} q(s_t \mid y_{1:t}) ds_t \\ &= \int \hat{w}_t(s_t) q(s_t \mid y_{1:t}) \, ds_t \\ &= \mathbb{E}_{q(s_t \mid y_{1:t})} \left[\hat{w}_t(s_t) \right] \\ \mathbb{E}_{p(s_t \mid y_{1:t})} \left[h(s_t) \right] &= \frac{1}{p(y_{1:t})} \int h(s_t) \frac{p(y_{1:t} \mid s_t) \, p(s_t)}{q(s_t \mid y_{1:t})} q(s_t \mid y_{1:t}) ds_t \\ &= \frac{\mathbb{E}_{q(s_t \mid y_{1:t})} \left[\hat{w}_t(s_t) h(s_t) \right]}{\mathbb{E}_{p(s_t \mid y_{1:t})} \left[\hat{w}_t(s_t) h(s_t) \right]} \end{aligned}$$

In practice

Draw $\left\{ \tilde{s}_{t}^{(i)} \right\}_{i=1:N}$ from the proposal $q(s_{t}|y_{1:t})$ and calculate their respective unormalized weights $\left\{ \hat{w}_{t}^{(i)} \right\}_{i=1:N}$.

$$\begin{split} \mathbb{E}_{p(s_t|y_{1:t})}\left[h(s_t)\right] &= \mathbb{E}_{q(s_t|y_{1:t})}\left[\tilde{w}_t(s_t)h(s_t)\right] \\ &= \frac{\mathbb{E}_{q(s_t|y_{1:t})}\left[\hat{w}_t(s_t)h(s_t)\right]}{\mathbb{E}_{q(s_t|y_{1:t})}\left[\hat{w}_t(s_t)\right]} \\ &\approx \frac{\frac{1}{N}\sum_{i=1}^N \hat{w}_t^{(i)}h(\tilde{s}_t^{(i)})}{\frac{1}{N}\sum_{i=1}^N \hat{w}_t^{(i)}} &= \sum_{i=1}^N \tilde{w}_t^{(i)}h(\tilde{s}_t^{(i)}) \end{split}$$

if we define the normalized weights as:

$$ilde{w}_t^{(i)} = rac{\hat{w}_t^{(i)}}{\sum_{i=1}^N \hat{w}_t^{(i)}}$$

Numerical integration (or perfect samplin Importance sampling Sequential Importance sampling Degeneracy problem and resampling Summary The likelihood expression

Sequential Importance sampling

A recursive approach can be implemented by choosing a *proposal* verifying:

$$q(s_t | y_{1:t}) = q(s_t | s_{t-1}, y_t) \ q(s_{t-1} | y_{1:t-1})$$

 s_t is simply drawn in $q(s_t|s_{t-1}, y_t)$.

The (unormalized) weights can be recursively calculated as:

$$\hat{w}_t(s_t) \propto \tilde{w}_{t-1}(s_{t-1}) \frac{p(y_t | s_t) p(s_t | s_{t-1})}{q(s_t | s_{t-1}, y_t)}$$

Skip Proof

Numerical integration (or perfect sampling Importance sampling Sequential Importance sampling Degeneracy problem and resampling Summary The likelihood expression

Unnormalized weights are defined by:

$$\begin{split} \hat{w}_{t}(s_{t}) &\equiv \frac{p(y_{1:t}|s_{t}) p(s_{t})}{q(s_{t}|y_{1:t})} \frac{p(y_{1:t-1}|s_{t-1}) p(s_{t-1})}{p(y_{1:t-1}|s_{t-1}) p(s_{t-1})} \\ &= \frac{p(y_{1:t}|s_{t}) p(s_{t})}{q(s_{t-1}|y_{1:t-1}) q(s_{t}|s_{t-1},y_{t})} \frac{p(y_{1:t-1}|s_{t-1}) p(s_{t-1})}{p(y_{1:t-1}|s_{t-1}) p(s_{t-1})} \\ &= \hat{w}_{t-1}(s_{t-1}) \frac{p(y_{1:t}|s_{t}) p(s_{t})}{p(y_{1:t-1}|s_{t-1}) p(s_{t-1}) q(s_{t}|s_{t-1},y_{t})} \end{split}$$

Since
$$p(s_t) = p(s_0) \prod_{i=1}^t p(s_i | s_{i-1})$$
 and $p(y_{1:t} | s_t) = \prod_{i=1}^t p(y_i | s_i)$:

$$\frac{p(y_{1:t}|s_t) p(s_t)}{p(y_{1:t-1}|s_{t-1}) p(s_{t-1})} = p(y_t|s_t) p(s_t|s_{t-1})$$

We obtain:

$$\hat{w}_{t}(s_{t}) = \hat{w}_{t-1}(s_{t-1}) \frac{p(y_{t}|s_{t})p(s_{t}|s_{t-1})}{q(s_{t}|s_{t-1},y_{t})}$$

$$\propto \tilde{w}_{t-1}(s_{t-1}) \frac{p(y_{t}|s_{t})p(s_{t}|s_{t-1})}{q(s_{t}|s_{t-1},y_{t})}$$

A generic particle filter

For t = 1, ..., T and i = 1, ..., N,

At time *t*, knowing $\left\{s_{t-1}^{(i)}, w_{t-1}^{(i)}\right\}_{i=1:N} (\approx p(s_{t-1}|y_{1:t-1}))$:

- Draw $\left\{\tilde{s}_{t}^{(i)}\right\}_{i=1:N}$ from $q\left(s_{t}|s_{t-1}^{(i)},y_{t}\right)$.
- $\quad \text{Evaluate the weights: } \hat{w}_t^{(i)} = w_{t-1}^{(i)} \frac{p(y_t | \tilde{s}_t^{(i)}) p(\tilde{s}_t^{(i)} | s_{t-1}^{(i)})}{q(\tilde{s}_t^{(i)} | s_{t-1}^{(i)}, y_t)}.$
- Use $\left\{ \tilde{\mathbf{s}}_t^{(i)}, \tilde{w}_t^{(i)} = \frac{\hat{w}_t^{(i)}}{\sum_{i=1}^N \hat{w}_t^{(i)}} \right\}_{i=1:N}$ as $\left\{ \mathbf{s}_t^{(i)}, w_t^{(i)} \right\}_{i=1:N}$ for the next filter iteration.

Numerical integration (or perfect sampling) Importance sampling Sequential Importance sampling Degeneracy problem and resampling Summary

Degeneracy problem and resampling

- No update here for the moment...
- As t increases, all-but-one particles have negligible weights (essentially in large samples).
- That is the reason why systematic resampling was initially proposed in the literature (Gordon et al., 1993).
- It consists in randomly drawing with replacement particles in their empirical distribution $\left\{\tilde{s}_{t}^{(i)}, \tilde{w}_{t}^{(i)}\right\}_{i=1:N}$.
- It amounts to discard particles with low weights and replicate particles with high weights to focus on interesting areas of the distribution using a constant number of particles.
- Negative side-effets
 - Impoverishment of the particles swarm.
 - It can render a filter iteration relatively time-consuming.
 - For estimation (see later).

Numerical integration (or perfect sampling) Importance sampling Sequential Importance sampling

Degeneracy problem and resampling Summary

The likelihood expression

Illustration

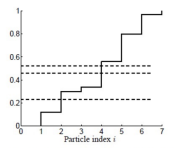


Figure 4.1: Illustrating the resampling step in the particle filter. The new set of particles is obtained by first generating M sorted uniformly distributed random numbers, three of which are shown by the dashed lines in the figure. These are then associated with a particle guided by the cumulative sum of the normalized importance weights. In the figure particle number 2 is chosen once and particle number 4 is chosen twice.

Sequential Importance-Sampling with Systematic Resampling

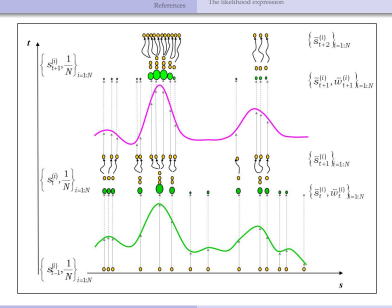
For t = 1, ..., T and i = 1, ..., N,

At time *t*, knowing $\left\{s_{t-1}^{(i)}, w_{t-1}^{(i)}\right\}_{i=1:N} (\approx p(s_{t-1}|y_{1:t-1}))$:

- Draw $\left\{\tilde{s}_{t}^{(i)}\right\}_{i=1:N}$ from $q\left(s_{t}|s_{t-1}^{(i)},y_{t}\right)$.
- $\quad \text{Evaluate the weights: } \hat{w}_t^{(i)} \propto w_{t-1}^{(i)} \frac{p(y_t | \mathbf{s}_t^{(i)}) p(\mathbf{s}_t^{(i)} | \mathbf{s}_{t-1}^{(i)})}{q(\mathbf{s}_t^{(i)} | \mathbf{s}_{t-1}^{(i)} y_t)}.$
- Resample

$$\left\{\tilde{s}_{t}^{(i)}, \tilde{w}_{t}^{(i)} = \frac{\hat{w}_{t}^{(i)}}{\sum_{i=1}^{N} \hat{w}_{t}^{(i)}}\right\}_{i=1:N} \mapsto \left\{s_{t}^{(i)}, w_{t}^{(i)} = \frac{1}{N}\right\}_{i=1:N} (\approx p(s_{t}|y_{1:t})).$$

Numerical integration (or perfect sampling Importance sampling Sequential Importance sampling Degeneracy problem and resampling Summary
The likelihood expression



Numerical integration (or perfect sampling)
Importance sampling
Sequential Importance sampling
Degeneracy problem and resampling
Summary
The likelihood expression

The sample likelihood

$$p(y_{1:T}|\Theta) = p(y_1|s_0;\Theta)p(s_0|\Theta) \prod_{t=2}^{T} p(y_t|y_{1:t-1};\Theta)$$

with

$$p(y_t | y_{1:t-1}; \Theta) \approx \sum_{i=1}^{N} \hat{w}_t^{(i)}$$

If $q(s_t|s_{t-1}, y_t; \Theta) = p(s_t|s_{t-1}; \Theta)$ and no pre-selection step:

$$p(y_t|y_{1:t-1};\Theta) \approx \sum_{i=1}^{N} \tilde{w}_{t-1}^{(i)} p(y_t|\tilde{s}_t^{(i)};\Theta)$$

In case of systematic resampling, since $\tilde{w}_{t-1}^{(i)} = 1/N$:

$$p(y_t|y_{1:t-1};\Theta) \approx \frac{1}{N} \sum_{i=1}^{N} p(y_t|\tilde{\mathbf{s}}_t^{(i)};\Theta)$$

Bayesian filtering methods Sequential Monte-Carlo methods The toy DSGE model The filters in Dynare

Conclusion

The model
Dynamic equilibrium
The steady state
True values
The .mod file

$$\mathbb{E}_0 \sum_{t=0}^{\infty} \beta^t u(c_t, l_t) = \mathbb{E}_0 \sum_{t=0}^{\infty} \beta^t \frac{\left(c_t^{\theta} (1 - l_t)^{1-\theta}\right)^{1-\tau}}{1-\tau}$$

subject to

$$\begin{aligned} y_t &= c_t + i_t \\ y_t &= a_t k_t^{\alpha} l_t^{1-\alpha} \\ k_{t+1} &= i_t + (1-\delta)k_t \\ log(a_{t+1}) &= \rho log(a_t) + \epsilon_t \end{aligned}$$

Skip deta

$$\frac{1-\theta}{\theta}\frac{c_t}{1-l_t} = (1-\alpha)\frac{y_t}{l_t}$$

$$\beta \mathbb{E}_t \left[\frac{\left(c_{t+1}^{\theta} (1-l_{t+1})^{1-\theta}\right)^{1-\tau}}{c_{t+1}} \left(1-\delta + \alpha \frac{y_{t+1}}{k_{t+1}}\right) \right] = \frac{\left(c_t^{\theta} (1-l_t)^{1-\theta}\right)^{1-\tau}}{c_t}$$

$$k_{t+1} = i_t + (1 - \delta)k_t$$

$$i_t = y_t - c_t$$

$$y_t = a_t k_t^{\alpha} l_t^{1-\alpha}$$

$$log(a_t) = \rho log(a_{t-1}) + \varepsilon_t$$

$$\bar{k} = \frac{-(\alpha - 1)\alpha^{\frac{1}{1 - \alpha}}\beta^{\frac{1}{1 - \alpha}}((\beta(\delta - 1) + 1)^{\frac{\alpha}{\alpha - 1}})\theta}{-\alpha\delta\beta + \delta\beta + \alpha\theta\beta - \beta - \alpha\theta + 1}$$

$$\bar{a} = 1$$

$$\overline{l} = \frac{(\alpha - 1)(\beta(\delta - 1) + 1)\theta}{\alpha\theta + \beta((\alpha - 1)\delta - \alpha\theta + 1) - 1}$$

References

$$\bar{y} = \bar{k}^{\alpha} \bar{l}^{1-\alpha}$$

$$\bar{i} = \delta \bar{k}$$

$$\bar{c} = \bar{y} - \bar{i}$$

The model Dynamic equilibrium The steady state **True values** The .mod file

Parameters		Benchmark	Risky	Extreme	Prior
Capital elasticity in the production function	α	0.4	0.4	0.4	$U_{[0;1]}$
Consumption-leisure substitution rate	θ	0.357	0.357	0.357	$U_{[0;1]}$
Discount factor	β	0.99	0.99	0.99	$U_{[0.75;1]}$
Depreciation rate of capital	δ	0.02	0.02	0.02	U _[0;0.05]
Persistence of technological shock	ρ	0.95	0.95	0.95	$U_{[0;1]}$
Intertemporal substitution	τ	2	50	50	$U_{[0;100]}$
Standard deviation of technological shock	σ_{ϵ}	0.007	0.035	0.035	$U_{[0;.1]}$
Standard deviation	σ_{v}	0.00158	0.0175	0.000158	$U_{[0;0.1]}$
of	σ_l	0.0011	0.00312	0.0011	$U_{[0;0.1]}$
measurement errors	σ_i	0.000866	0.00465	0.000866	U _[0;0.1]
Datafile:		benchmark	risky	extreme	

Declaration and assignment

```
var k A c l i y;
varexo e_a;

parameters alp bet tet tau delt rho;

alp = 0.4;
bet = 0.99;
tet = 0.357;
tau = 50;
delt = 0.02;
rho = 0.95;
```

Model declaration

```
model:
c = ((1 - alp) *tet/(1-tet)) *A*(1-1) *((k(-1)/1)^alp);
y = A*(k(-1)^alp)*(l^(1-alp));
i = v-c;
k = (1-delt) * k(-1) + i ;
log(A) = rho*log(A(-1)) + e_a;
(((c^{(tet)})*((1-1)^{(1-tet)}))^{(1-tau)})/c -
\text{bet}*((((c(+1)^(tet))*((1-1(+1))^(1-tet)))^(1-tau))/c(+1))*
(1 - delt + alp * (A(1) * (k^alp) * (l(1)^(1-alp)))/k) = 0;
end:
shocks:
var e a; stderr 0.035;
end:
steady;
```

The model Dynamic equilibrium The steady state True values The .mod file

Measurement errors

Be careful: the measurement errors should not be declared in the block shocks!

The model
Dynamic equilibrium
The steady state
True values
The .mod file

Priors

```
estimated_params;
alp, uniform_pdf,,, 0.0001, 1;
bet, uniform_pdf,,, 0.75, 0.999;
tet, uniform_pdf,,, 0.0001, 1;
tau, uniform_pdf,,, 0.0001, 100;
delt, uniform_pdf,,, 0.0001, 0.05;
rho, uniform_pdf,,, 0.0001, 0.999;
stderr e_a, uniform_pdf,,, 0.00001, 0.1;
stderr y, uniform_pdf,,, 0.00001, 0.1;
stderr l, uniform_pdf,,, 0.00001, 0.1;
stderr i, uniform_pdf,,, 0.00001, 0.1;
end;
```

The model
Dynamic equilibrium
The steady state
True values
The .mod file

Starting values

```
estimated_params_init;
alp, 0.4;
bet, 0.99;
tet, 0.357;
tau, 50;
delt, 0.02;
rho, 0.95;
stderr e_a, .035;
stderr v, .0175;
stderr 1, .00312;
stderr i, .00465;
end;
varobs y l i;
```

For simplicity sake, a usual choice for the proposal is:

$$q(s_t|s_{t-1},y_t) = p(s_t|s_{t-1})$$

The weight expression simplifies:

$$\hat{w}_{t}(s_{t}) \propto \tilde{w}_{t-1}(s_{t-1}) \frac{p(y_{t} | s_{t}) p(s_{t} | s_{t-1})}{q(s_{t} | s_{t-1}, y_{t})}$$

$$= \tilde{w}_{t-1}(s_{t-1}) p(y_{t} | s_{t})$$

Easy to write since measurement errors are assumed additive and gaussian (remember $y_t = g(s_t; \Theta) + e_t$):

$$p(y_t | \tilde{\mathbf{g}}_t^{(i)}) = (2\pi)^{-\frac{\dim(y_t)}{2}} |P_e|^{-\frac{1}{2}} \exp \left\{ -\frac{1}{2} \left[y_t - g(\tilde{\mathbf{g}}^{(i)}; \Theta) \right]' P_e^{-1} \left[y_t - g(\tilde{\mathbf{g}}^{(i)}; \Theta) \right] \right\}$$

sometimes difficult to calculate due to $V(e_t) \equiv P_e$.

The SISR particle filter

The Auxiliary particle filter Gaussian approximations The nonlinear Kalman filter Marginal (particle) filters Conditional particle filters

with Dynare

```
estimation( order= 2,

number_of_particles= [5000],

resampling = [systematic], none, generic,

resampling_method = [kitagawa], residual, smooth,

mh_replic = 0,

mode_compute= 7 or 8);
```

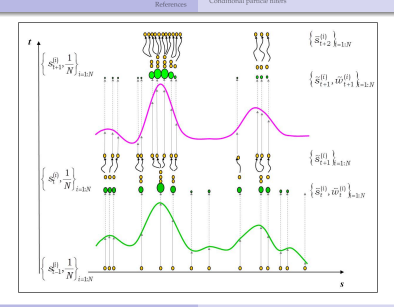
The SISR particle filter
The Auxiliary particle filter
Gaussian approximations
The nonlinear Kalman filter
Marginal (particle) filters
Conditional particle filters

- Can be viewed as a way to reintroduce y_t in the proposal.
- Add a pre-selection (resampling) step on the past particles based on the predictive likelihood when it is more informative than the transition distribution.
- Only the most promising particles will enter in the calculation of the current particle *proposal* distribution.
- Implement the previous filter on pre-selected past particles (modify slightly current particle weights).
- The resampling step on current particles becomes optional.

Bayesian filtering methods Sequential Monte-Carlo methods The toy DSGE model The filters in Dynare

Conclusion

The SISR particle filter
The Auxiliary particle filter
Gaussian approximations
The nonlinear Kalman filter
Marginal (particle) filters



- Suppose we know $\left\{s_{t-1}^{(i)}, w_{t-1}^{(i)}\right\}_{i=1:N} (\approx p(s_{t-1}|y_{1:t-1})).$
- Approximate ε_t with $\left\{\chi_j^{\varepsilon_t}, W_j\right\}_{j=1:L}$ and the state predictive density such as: $\bar{s}_t^{(i)} = \int f(s_{t-1}^{(i)}, \varepsilon_t; \Theta) N(\varepsilon_t; 0, Q) d\varepsilon_t \approx \sum_{j=1}^L W_j f(s_{t-1}^{(i)}, \chi_j^{\varepsilon_t}; \Theta)$ or simply set $\bar{s}_t^{(i)} = f(s_{t-1}^{(i)}, 0; \Theta)$
- Define $\hat{\tau}_{t-1}^{(i)} \equiv p(y_t | \bar{s}_t^{(i)}) \ w_{t-1}^{(i)}$ and resample $\left\{ s_{t-1}^{(i)}, \tilde{\tau}_{t-1}^{(i)} = \frac{\hat{\tau}_{t-1}^{(i)}}{\sum_{i=1}^{N} \hat{\tau}_{t-1}^{(i)}} \right\}_{i=1:N}$ $\mapsto \left\{ s_{t-1}^{(k)}, \frac{1}{N} \right\}_{1:N}$.
- Draw $\left\{\tilde{s}_{t}^{(l)}\right\}_{l=1:N}$ from $q\left(s_{t}|s_{t-1}^{(k^{l})},y_{t}\right)$.
- $\begin{array}{l} \bullet \text{ Evaluate the weights: } \hat{w}_t^{(l)} \propto \underbrace{\frac{w_{t-1}^{(k')}}{\tilde{\tau}_{t-1}^{(k')}}}_{\text{stage 1 weights}} \underbrace{\frac{1}{N} \frac{p(y_t | \tilde{\mathbf{s}}_t^{(l)}) p(\tilde{\mathbf{s}}_t^{(l)} | \mathbf{s}_{t-1}^{(k')})}{q(\tilde{\mathbf{s}}_t^{(l)} | \mathbf{s}_{t-1}^{(k')}, y_t)}}_{\text{stage 2 weights}}. \end{array}$
- We get $\left\{ \tilde{s}_t^{(l)}, \tilde{w}_t^{(l)} = \frac{\hat{w}_t^{(l)}}{\sum_{l=1}^{l} \hat{w}_t^{(l)}} \right\}_{l=1:N} = \left\{ s_t^{(l)}, w_t^{(l)} \right\}_{l=1:N}$ (no resampling).

• If $q(s_t|s_{t-1}, y_t) = p(s_t|s_{t-1})$, the weight becomes

$$\hat{w}_{t}^{(l)} \propto \frac{1}{N} \frac{p(y_{t} | \tilde{\mathbf{s}}_{t}^{(l)})}{p(y_{t} | \tilde{\mathbf{s}}_{t}^{(k')})} \sum_{i=1}^{N} \hat{\boldsymbol{\tau}}_{t-1}^{(l)}$$

• Optional resampling in $\left\{ \tilde{s}_t^{(l)}, \tilde{w}_t^{(l)} = \frac{\hat{w}_t^{(l)}}{\sum_{l=1}^N \hat{w}_t^{(l)}} \right\}_{l=1:N} => \text{the SIR filter with a pre-selection step.}$

The SISR particle filter
The Auxiliary particle filter
Gaussian approximations
The nonlinear Kalman filter
Marginal (particle) filters
Conditional particle filters

with Dynare

```
estimation( order= 2,

filter_algorithm= apf,

number_of_particles= [5000],

resampling = [systematic],none,

mh_replic = 0,

mode_compute= 7 or 8);
```

The SISR particle filter
The Auxiliary particle filter
Gaussian approximations
The nonlinear Kalman filter
Marginal (particle) filters
Conditional particle filters

	DGP	Kalman	SIR	APF
α	0.400	0.4046	0.4003	0.4007
		(0.0032)		
θ	0.357	0.3566	0.3574	0.3574
		(0.0013)		
β	0.990	0.9977	0.9891	0.9890
		(0.0000)		
δ	0.020	0.0215	0.0199	0.0199
		(0.0007)		
ρ	0.950	0.9743	0.9508	0.9491
		(0.0017)		
τ	50.000	54.3064	49.9102	49.9216
	0.005	(11.9222)	0.0055	0.0045
σ ε	0.035	0.0438	0.0355	0.0365
_	0.0175	(0.0026)	0.0181	0.0179
σ_y	0.0175	0.000	0.0181	0.0179
-	0.00312	(0.0012) 0.0037	0.0310	0.0310
σ_l	0.00312	(0.0002)	0.0310	0.0310
<i>a</i> .	0.00465	0.0002	0.0047	0.0046
σ_i	0.00403	(0.0000)	0.0047	0.0040
		(0.0000)		
Posterior Kernel		1,155,302	1.187.612	1.187.221
Proposal approximation		1,133.302	Particles	Particles
Distribution approximation			Particles	Particles
N			10,000	10,000
Mixture		_	-	-
Option		order=1	order=2	order=2
Resampling		-	systematic	none
			.,	
				l

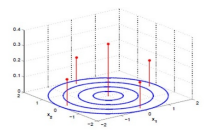
The SISR particle filter The Auxiliary particle filter Gaussian approximations The nonlinear Kalman filter Marginal (particle) filters Conditional particle filters

Gaussian approximations

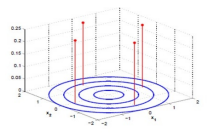
- The distribution of particles is assumed to be Gaussian.
- Built with Monte-Carlo approximations or Gaussian sparse grids.
 - the Cubature points (Arasaratnam and Haykin, 2009)
 - the Unscented sigma-points (Julier and Ulhmann, 1997)
 - the Smolyak quadrature (Winschel and Kratzig, 2010)
- Pros: fast and easy to implement
 - drastically reduces the number of operations wrt Monte Carlo.
 - focus on the mean and variance of a unique Gaussian distribution
 - no resampling necessary (in theory)
- Cons: extreme approximation, highly inaccurate in case of multimodal distributions.

The SISR particle filter
The Auxiliary particle filter
Gaussian approximations
The nonlinear Kalman filter
Marginal (particle) filters
Conditional particle filters

Illustration in a bivariate case



(a) Sigma point set for the UKF



(b) Third-degree spherical-radial cubature point set for the CKF

- Insert the nonlinear model in the Kalman filter: the moments of all gaussian densities are numerically approximated.
- Suppose the state distribution is Gaussian so that we only track $\bar{s}_{t|t}$ and $P_{s_{t|t}}$.
- Suppose we know $N(s_{t-1}; \bar{s}_{t-1|t-1}, P_{s_{t-1|t-1}})$ at date t.
- Approximate (s_{t-1}, ε_t) with sigma-points $\{\chi_j, W_j\}_{j=1:L}$ where $\chi_j = \left(\chi_j^{s_{t-1}}, \chi_j^{\varepsilon}\right)$.
- Deduce the densities of s_t and y_t using the (nonlinear) state-space representation:

$$\chi_j^{s_t} = f(\chi_j^{s_{t-1}}, \chi_j^{\varepsilon}; \Theta)$$

$$\chi_j^{y_t} = g(\chi_j^{s_t}; \Theta)$$

Empirical moments computation

$$\begin{split} \bar{\mathbf{s}}_{t|t-1} &= \sum_{j=1}^{L} W_{j} \, \chi_{j}^{s_{t}} \\ P_{s_{t|t-1}} &= \sum_{j=1}^{L} W_{j} \left(\chi_{j}^{s_{t}} - \bar{\mathbf{s}}_{t|t-1} \right) \left(\chi_{j}^{s_{t}} - \bar{\mathbf{s}}_{t|t-1} \right)^{'} \\ \bar{y}_{t|t-1} &= \sum_{j=1}^{L} W_{j} \, \chi_{j}^{y_{t}} \\ P_{y_{t|t-1}} &= \sum_{j=1}^{L} W_{j} \left(\chi_{j}^{y_{t}} - \bar{y}_{t|t-1} \right) \left(\chi_{j}^{y_{t}} - \bar{y}_{t|t-1} \right)^{'} + P_{e} \\ P_{s_{t|t-1}, y_{t|t-1}} &= \sum_{j=1}^{L} W_{j} \left(\chi_{j}^{s_{t}} - \bar{\mathbf{s}}_{t|t-1} \right) \left(\chi_{j}^{y_{t}} - \bar{y}_{t|t-1} \right)^{'} \end{split}$$

Kalman update step

• Calculate the Kalman gain $K_t = P_{s_{t|t-1}, y_{t|t-1}} P_{y_{t|t-1}}^{-1}$ and deduce the *posterior* moments of state variables:

$$\begin{split} \bar{s}_{t|t} &= \bar{s}_{t|t-1} + K_{t} \left(y_{t} - \bar{y}_{t|t-1} \right) \\ P_{s_{t|t}} &= P_{s_{t|t-1}} - K_{t} P_{y_{t|t-1}} K_{t}^{'} \end{split}$$

- Of course imperfect since the Kalman updating step derived for a gaussian/linear framework is implemented in a nonlinear framework.
- Approximative likelihood: $p(y_t|y_{1:t-1};\Theta) \approx N(y_t; \bar{y}_{t|t-1}, P_{y_{t|t-1}})$.

The SISR particle filter The Auxiliary particle filter Gaussian approximations The nonlinear Kalman filter Marginal (particle) filters Conditional particle filters

with Dynare

```
estimation( order= 2,
filter_algorithm= nlkf,
proposal_approximation= cubature, [unscented], montecarlo,
mh_replic = 0,
mode_compute= );
```

The SISR particle filter The Auxiliary particle filter Gaussian approximations The nonlinear Kalman filter Marginal (particle) filters Conditional particle filters

	DGP		NLKF	
α	0.400	0.3411	0.3390	0.3407
θ	0.357	0.3344	0.3334	0.3343
β	0.990	0.9956	0.9957	0.9955
δ	0.020	0.0101	0.0095	0.0101
ρ	0.950	0.9786	0.9793	0.9785
τ	50.000	20.3817	21.6798	20.3379
σ_{ϵ}	0.035	0.0410	0.0413	0.0411
σ_y	0.0175	0.0178	0.0176	0.0177
σ_l	0.00312	0.0031	0.0031	0.0031
σ_i	0.00465	0.0000	0.0000	0.0001
Posterior Kernel Proposal approximation Distribution approximation N Mixture Option Resampling		1,200.734 - Particles 10,000 - -	1,200.804 	1,200.249

References

The SISR particle filter The Auxiliary particle filter Gaussian approximations The nonlinear Kalman filter Marginal (particle) filters Conditional particle filters

Intuition

- Suppose a gaussian-based distribution for the states.
- Build the nonlinear Kalman filter posterior.
- Build an approximation for the *proposal*: adds an extra layer of temporary particles (or sparse grid) with weights wrt the nonlinear Kalman filter.
- Two approximations here: gaussian and gaussian-mixture (more general).

Gaussian (particle) filters

- Suppose we know $N(s_{t-1}; \tilde{s}_{t-1}, P_{s_{t-1}})$ at date t.
- Build $N(s_t; \bar{s}_{t|t-1}, P_{s_{t|t-1}})$ and $N(s_t; \bar{s}_{t|t}, P_{s_{t|t}})$, resp. the *prior* and *posterior* distributions using a sparse-grid nonlinear Kalman filter.
- Draw $\left\{\tilde{s}_{t}^{(i)}\right\}_{i=1:N}$ in $N(s_{t}; \bar{s}_{t|t}, P_{s_{t|t}})$.
- Calculate their weights: $\hat{w}_t^{(i)} = \frac{1}{N} \frac{p(y_t | \bar{s}_t^{(i)}) N(\bar{s}_t^{(i)}; \bar{s}_{t|t-1}, P_{\bar{s}_{t|t-1}})}{N(\bar{s}_t^{(i)}; \bar{s}_{t|t}, P_{\bar{s}_{t|t}})}$
- Compute the moments from particles (eventually resampled):

$$\tilde{s}_t = \sum_{i=1}^{N} \tilde{w}_t^{(i)} \tilde{s}_t^{(i)}$$
 and $P_{s_t} = \sum_{i=1}^{N} \tilde{w}_t^{(i)} (\tilde{s}_t^{(i)} - \tilde{s}_t) (\tilde{s}_t^{(i)} - \tilde{s}_t)'$

The SISR particle filter
The Auxiliary particle filter
Gaussian approximations
The nonlinear Kalman filter
Marginal (particle) filters
Conditional particle filters

with Dynare

```
estimation(
                                   order=
                         filter_algorithm=
                                            gf,
                proposal_approximation=
                                            cubature, [unscented],
             distribution_approximation=
                                            cubature, [unscented], montecarlo,
                    number_of_particles=
                                            [5000],
                             resampling =
                                            [systematic], none, generic,
                    resampling_method =
                                            [kitagawa], residual, smooth,
                              mh_replic =
                                            0.
                         mode_compute=
                                            depends if resampling or not);
```

The SISR particle filter
The Auxiliary particle filter
Gaussian approximations
The nonlinear Kalman filter
Marginal (particle) filters
Conditional particle filters

	DGP			GF		
α	0.400	0.3995	0.3990	0.3615	0.3580	0.3503
θ	0.357	0.3559	0.3557	0.3409	0.3394	0.3363
β	0.990	0.9901	0.9904	0.9946	0.9952	0.9966
δ	0.020	0.0195	0.0194	0.0119	0.0109	0.0095
ρ	0.950	0.9581	0.9590	0.9743	0.9764	0.9802
τ	50.000	37.8044	37.8781	30.9630	33.9064	28.8161
σε	0.035	0.0422	0.0413	0.0418	0.0396	0.0418
σ_y	0.0175	0.0178	0.0179	0.0176	0.0170	0.0171
σ_l	0.00312	0.0031	0.0031	0.0031	0.0031	0.0031
σ_i	0.00465	0.0067	0.0065	0.0058	0.0083	0.0075
Posterior Kernel Proposal approximation Distribution approximation N Mixture Option Resampling		1,180.992 Unscented Particles 10,000	1,181.274 Particles Particles 10,000 - systematic	1,189.404 Unscented Particles 10,000	1,174.907 Unscented Unscented - - -	1,176.602 Cubature Cubature

Gaussian-mixture (particle) filters

- Kotesha and Djuric (2003), van der Merwe and Wan (2009)
- Limits the size of the Gaussian-mixture for states.
- A *G*-Gaussian mixture $\{\alpha^{(g)}, \mu^{(g)}, P^{(g)}\}_{g=1:G}$ is *G* distributions $N(s; \mu^{(g)}, P^{(g)})$ combined with weights $\alpha^{(g)}$ (with $\sum_{g=1}^{G} \alpha^{(g)} = 1$).

The SISR particle filter
The Auxiliary particle filter
Gaussian approximations
The nonlinear Kalman filter
Marginal (particle) filters
Conditional particle filters

Suppose we have

$$p_{GM(G)}(s_{t-1}|y_{1:t-1}) = \sum_{g=1}^{G} \alpha_{t-1}^{(g)} N(s_{t-1}; \mu_{t-1}^{(g)}, P_{t-1}^{(g)})$$

$$p_{GM(H)}(\varepsilon_t) = \sum_{h=1}^{H} \beta_t^{(h)} N(\varepsilon_t; \mu_{\varepsilon_t}^{(h)}, Q_t^{(h)})$$

References

We track $\{\alpha_t^{(g)}, \mu_t^{(g)}, P_t^{(g)}\}$.

- Define g'(=1,...,G'=GH).
- The nonlinear Kalman filter is implemented on each combinated element of the mixtures to build the *prior* and *posterior* Gaussian-mixture approximations:

$$p_{GM(G')}(s_t | y_{1:t-1}) = \sum_{g'=1}^{G'} \alpha_t^{(g')} N(s_t; \mu_t^{(g')}, P_t^{(g')})$$

$$p_{GM(G')}(s_t | y_{1:t}) = \sum_{g''=1}^{G'} \alpha_t^{(g'')} N(s_t; \mu_t^{(g'')}, P_t^{(g'')})$$

• The *posterior* Gaussian-mixture $p_{GM(G')}(s_t|y_{1:t})$ is used as proposal.

- Draw current particles $\left\{ \tilde{s}_{t}^{(i)} \right\}_{i=1:N}$ in $p_{GM(G')}(s_{t}|y_{1:t})$
- The weight $\hat{w}_t^{(i)}$ is provided by:

$$\hat{w}_{t}^{(i)} = \frac{1}{N} \frac{p(y_{t} | \tilde{\mathbf{S}}_{t}^{(i)}) \ p_{GM(G')}(\tilde{\mathbf{S}}_{t}^{(i)} | y_{1:t-1})}{p_{GM(G')}(\tilde{\mathbf{S}}_{t}^{(i)} | y_{1:t})}$$

• Enventually resample and fit a new *G*-Gaussian mixture $\left\{\alpha^{(g)}, \mu^{(g)}, P^{(g)}\right\}_{g=1:G} \text{ on particles } \left\{\tilde{s}_t^{(i)}, \tilde{w}_t^{(i)} = \frac{\hat{w}_t^{(i)}}{\sum_{i=1}^N \hat{w}_t^{(i)}}\right\}_{i=1:N} \text{ to avoid explosion.}$

The SISR particle filter
The Auxiliary particle filter
Gaussian approximations
The nonlinear Kalman filter
Marginal (particle) filters
Conditional particle filters

with Dynare

```
estimation(
                                   order=
                         filter_algorithm=
                                            gmf,
                proposal_approximation=
                                            cubature,[unscented],
             distribution_approximation=
                                            cubature, [unscented], montecarlo,
                    number_of_particles=
                                            [5000],
                             resampling =
                                            [systematic], none, generic,
                    resampling_method =
                                            [kitagawa], residual, smooth,
                              mh_replic =
                                            0,
                         mode_compute=
                                            depends if resampling or not);
```

The SISR particle filter
The Auxiliary particle filter
Gaussian approximations
The nonlinear Kalman filter
Marginal (particle) filters
Conditional particle filters

To be completed...

	DGP	GM	IF
α	0.400		0.3812
θ	0.357		0.3487
β	0.990		0.9936
δ	0.020		0.0154
ρ	0.950		0.9703
τ	50.000		47.1903
σ_{ϵ}	0.035		0.0305
σ_y	0.0175		0.0172
σ_l	0.00312		0.0031
σ_i	0.00465		0.0072
Posterior Kernel Proposal approximation Distribution approximation N Mixture Option Resampling		1,189.1608 Unscented Particles 10,000 5 - systematic	1,182.237 Unscented Unscented - 5 -

The SISR particle filter The Auxiliary particle filter Gaussian approximations The nonlinear Kalman filter Marginal (particle) filters Conditional particle filters

Intuition

- No particular distributional assumption on states.
- Combine each particle with a Gaussian approximation on structural shocks.
- Still use the nonlinear Kalman filter *posterior* as *proposal*, but for each particle.

The SISR particle filter The Auxiliary particle filter Gaussian approximations The nonlinear Kalman filter Marginal (particle) filters Conditional particle filters

- Suppose we know $\left\{s_{t-1}^{(i)}, w_{t-1}^{(i)}\right\}_{i=1:N}$.
- Approximate ε_t with sigma-points $\left\{\chi_j^{\varepsilon}, W_j\right\}_{j=1:L}$:

$$\begin{aligned} \chi_j^{s_t^{(i)}} &= f(s_{t-1}^{(i)}, \chi_j^{\varepsilon}; \Theta) \\ \chi_j^{y_t^{(i)}} &= g(\chi_j^{s_t^{(i)}}; \Theta) \end{aligned}$$

- Proposal: draw $\left\{\tilde{s}_{t}^{(i)}\right\}_{i=1:N}$ in $N(s_{t}; \bar{s}_{t|t}^{(i)}, P_{s_{t|t}}^{(i)})$, the Kalman *posterior*.
- Approximated weights:

$$\hat{w}_t^{(i)} pprox w_{t-1}^{(i)} N(y_t; \bar{y}_{t|t-1}^{(i)}, P_{y_{t|t-1}}^{(i)})$$
 a la Amisano and Tristani (2010)

$$\hat{w}_{t}^{(i)} \approx w_{t-1}^{(i)} \frac{p(y_{t}|\tilde{\mathbf{s}}_{t}^{(i)})N(\tilde{\mathbf{s}}_{t}^{(i)};\tilde{\mathbf{s}}_{t|t-1}^{(i)}, P_{\tilde{\mathbf{s}}_{t|t-1}}^{(i)})}{N(\tilde{\mathbf{s}}_{t}^{(i)};\tilde{\mathbf{s}}_{t|t}^{(i)}, P_{\tilde{\mathbf{s}}_{t|t}}^{(i)})} \ a \ la \ \text{Murray et al. (2013)}$$

• Resample
$$\left\{ \tilde{s}_t^{(i)}, \tilde{w}_t^{(i)} = \frac{\hat{w}_t^{(i)}}{\sum_{l=1}^N \hat{w}_t^{(i)}} \right\}_{i=1:N} \mapsto \left\{ s_t^{(i)}, w_t^{(i)} = \frac{1}{N} \right\}_{i=1:N}$$
.

The SISR particle filter
The Auxiliary particle filter
Gaussian approximations
The nonlinear Kalman filter
Marginal (particle) filters
Conditional particle filters

Calling estimation

```
estimation( order= 2,
    filter_algorithm= cpf,
    proposal_approximation= cubature,[unscented],
    number_of_particles= [5000],
    cpf_weights = [amisanotristani], murrayjonesparslow,
    mh_replic = 0,
    mode_compute= 7 or 8);
```

The SISR particle filter
The Auxiliary particle filter
Gaussian approximations
The nonlinear Kalman filter
Marginal (particle) filters
Conditional particle filters

	DGP		CPF
α	0.400	0.4009	0.4026
θ	0.357	0.3566	0.3570
β	0.990	0.9896	0.9905
δ	0.020	0.0194	0.0198
ρ	0.950	0.9562	0.9594
τ	50.000	46.6954	48.4266
σ_{ϵ}	0.035	0.0395	0.0354
σ_y	0.0175	0.0183	0.0180
σ_l	0.00312	0.0031	0.0030
σ_i	0.00465	0.0038	0.0057
Posterior Kernel Proposal approximation Distribution approximation N Mixture cpf_weights Resampling		1,191.514 Unscented Particles 1,000 - amisanotristani systematic	1,176.0563 Unscented Particles 1,000 murrayjonesparslow systematic

The likelihood expression
The limits of the likelihood approach
Particle MCMC
The online approach

The sample likelihood

$$p(y_{1:T}|\Theta) = p(y_1|s_0;\Theta)p(s_0|\Theta) \prod_{t=2}^{T} p(y_t|y_{1:t-1};\Theta)$$

with

$$p(y_t | y_{1:t-1}; \Theta) \approx \sum_{i=1}^{N} \hat{w}_t^{(i)}$$

If $q(s_t|s_{t-1}, y_t; \Theta) = p(s_t|s_{t-1}; \Theta)$ and no pre-selection step:

$$p(y_t|y_{1:t-1};\Theta) \approx \sum_{i=1}^{N} \tilde{w}_{t-1}^{(i)} p(y_t|\tilde{s}_t^{(i)};\Theta)$$

In case of systematic resampling, since $\tilde{w}_{t-1}^{(i)} = 1/N$:

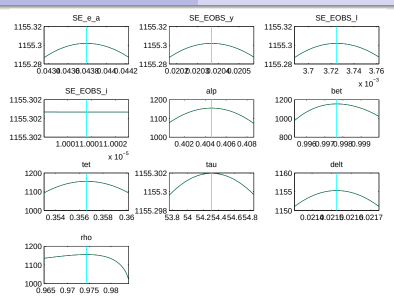
$$p(y_t|y_{1:t-1};\Theta) \approx \frac{1}{N} \sum_{i=1}^{N} p(y_t|\tilde{\mathbf{s}}_t^{(i)};\Theta)$$

The likelihood expression
The limits of the likelihood approach
Particle MCMC
The online approach

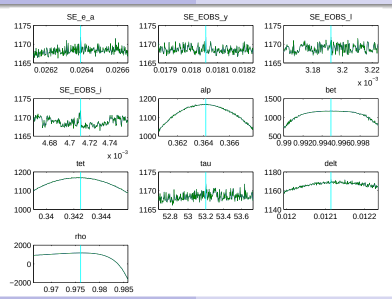
The limits of the likelihood approach

- Resampling is necessary to avoid degeneracy...
- ... but complicates inference through maximum likelihood because it renders the likelihood criterion nonsmooth.
- Even when the seed for random draws is fixed across the simulations, the traditional likelihood estimator depends on both resampled particles and the unknown parameters. A small change in the parameters value will cause a small change in the importance weights that will potentially generate a different set of resampled particles. This produces a discontinuity in the likelihood criterion.

The likelihood expression The limits of the likelihood approach Particle MCMC The online approach



The likelihood expression The limits of the likelihood approach Particle MCMC The online approach



The likelihood expression The limits of the likelihood approach Particle MCMC The online approach

Solutions

- No resample and implement maximum likelihood if possible.
- Resample and implement maximum likelihood with no gradient-based maximization method (downhill simplex, S.A., CMAES, ...).
- Implement a MCMC approach to build the posterior distribution of parameters (Andrieu et al. (2010)).
- Implement an online approach, i.e. consider parameters as extra states.
- Use a smooth resampling method (but requires extensions to multivariate state variables).
- SMC methods: coming soon

Particle Marginal Metropolis-Hastings (P-MMH)

- Build $p(\Theta | y_{1:T})$, the *posterior* distribution of parameters Θ .
- As previously, it is a function of the sample likelihood $p(y_{1:T}|\Theta)$ and the *priors* on parameters $p(\Theta)$:

$$p(\Theta|y_{1:T}) \propto p(y_{1:T}|\Theta)p(\Theta)$$

- In the case of a linear model, the sample likelihood $p(y_{1:T} | \Theta)$ is provided by the Kalman filter.
- In the case of a nonlinear model, an unbiased approximation of the sample likelihood $p(y_{1:T}|\Theta)$ is provided by particle filtering (Delmoral, 2004).

Particle Random-Walk Metropolis algorithm (Andrieu et al., 2010)

• For j = 1, ..., M (set large), define a candidate as:

$$\Theta_{j}^{\star} = \Theta_{j-1} + \epsilon_{j}$$

with $\epsilon_j \sim N(0, \gamma_{RW}V(\Theta_0))$ and γ_{RW} set in order to obtain an acceptance ratio around 24%.

 The posterior distribution of deep parameters can be approximated thanks to the acceptation rule:

$$\Theta_{j} = \begin{cases} \Theta_{j}^{\star} & \text{if } U_{[0,1]} \leqslant \min\left\{1, \frac{p(\Theta_{j}^{\star}|y_{1:T}|)}{p(\Theta_{j-1}|y_{1:T}|)}\right\} \\ \Theta_{j-1} & \text{otherwise} \end{cases}$$

The likelihood expression
The limits of the likelihood approach
Particle MCMC
The online approach

with Dynare

estimation(order= 2. filter_algorithm= [sis], nlkf, gf, gmf, apf, cpf, proposal_approximation= cubature, [unscented], distribution_approximation= cubature, [unscented], montecarlo, number_of_particles= [5000], resampling = [systematic], none, generic, resampling_method = [kitagawa], residual, smooth, mh_replic = [20000], mode_file= xx)

The likelihood expression The limits of the likelihood approach Particle MCMC The online approach

The online approach (Liu and West, 2001)

- Unknown parameters are considered as extra state variables to allow *online* evaluation. Combines the auxiliary particle filter with an assumed importance density $p(\Theta_t | \Theta_{t-1})$ for parameters.
- Produces time-varying parameters and thus adds noise to the parameter estimates. To reduce the effect of the artificial variability, the authors adopt a kernel shrinkage technique.
- Pros:
 - Requires only one pass over the sample.
 - Also works if order=1.
- Cons:
 - Two resampling steps (on past and current particles)
 - No strong theoretical foundations but recent developments.

The likelihood expression The limits of the likelihood approach Particle MCMC

The online approach

Suppose we have $\left\{ s_{t-1}^{(i)}, \Theta_{t-1}^{(i)}, w_{t-1}^{(i)} \right\}_{i=1:N}$. Calculate:

$$\begin{split} &\bar{\Theta}_{t-1} = \sum_{i=1}^{N} w_{t-1}^{(i)} \Theta_{t-1}^{(i)} \\ &m_{t-1}^{(i)} = a \, \Theta_{t-1}^{(i)} + (1-a) \, \bar{\Theta}_{t-1} \\ &V_{t-1} = \sum_{i=1}^{N} w_{t-1}^{(i)} (\Theta_{t-1}^{(i)} - \bar{\Theta}_{t-1}) (\Theta_{t-1}^{(i)} - \bar{\Theta}_{t-1})' \\ &\bar{s}_{t}^{(i)} = f(s_{t-1}^{(i)}, 0; m_{t-1}^{(i)}) \end{split}$$

- The index k^l is obtained from sampling in $\hat{\tau}_{t-1}^{(i)} \propto w_{t-1}^{(i)} p(y_t | \bar{s}_t^{(i)}, m_{t-1}^{(i)})$.

- **1** The weights are calculated as: $\hat{w}_t^{(l)} \propto p(y_t | \tilde{\mathbf{s}}_t^{(l)}, \tilde{\boldsymbol{\Theta}}_t^{(l)}) \frac{w_{t-1}^{(k^l)}}{\tilde{\tau}_{t-1}^{(k^l)}}$ and normalized.

- The shrinkage technique (based on parameter *a*) is used to produce slowly time-varying parameters and also to limit the variability.
- δ is the key parameter that conditions the shrinkage and the smoothness parameters a and b:

$$a = \frac{3\delta - 1}{2\delta}$$
$$b^2 = 1 - a^2$$

• δ is generally chosen in the range [0.95; 0.99].

The likelihood expression The limits of the likelihood approach Particle MCMC The online approach

with Dynare

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\begin{array}{ll} estimation( & order = & 1 \ or \ 2, \\ number\_of\_particles = & [5000] \ , \\ resampling = & [systematic], \ none, \\ mode\_compute = & 11) \ ; \\ \\ and \ fix \ \delta \ with \ options\_particle. \\ \hline liu\_west\_delta = 0.9 \end{array}
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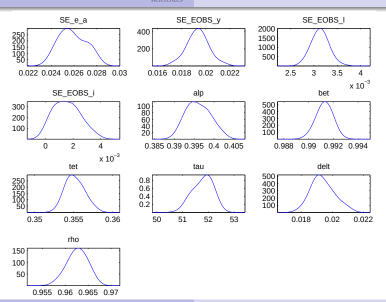
The likelihood expression The limits of the likelihood approach Particle MCMC

The online approach

	DGP	P-MMH	Online	
α	0.400	0.3960	0.42843	0.43621
	İ	[0.3908; 0.4003]	[0.39500.4621]	[0.4071; 0.4852]
θ	0.357	0.3551	0.36648	0.36603
		[0.3533; 0.3571]	[0.3004; 0.4193]	[0.3281; 0.4474]
β	0.990	0.9913	0.96926	0.97778
		[0.9902; 0.9923]	[0.9278; 0.9963]	[0.9178; 0.9982]
δ	0.020	0.0194	0.02473	0.02481
		[0.0181; 0.0205]	[0.0184; 0.0323]	[0.0176; 0.0345]
ρ	0.950	0.9626	0.67283	0.75078
	İ	[0.9590; 0.9662]	[0.5210; 0.8282]	[0.5762; 0.8620]
τ	50.000	51.7763	40.37328	66.82140
	İ	[51.1611; 52.3328]	[20.7203;54.2846]	[30.5316; 94.5316]
σ ε	0.035	0.0259	0.01825	0.02783
		[0.0242; 0.0277]	[0.0023; 0.0340]	[0.0095; 0.0403]
$\sigma_{\mathcal{U}}$	0.0175	0.0193	0.07411	0.05547
Ÿ		[0.0175; 0.0206]	[0.0510; 0.0949]	[0.0364; 0.0735]
σ_{l}	0.00312	0.0031	0.03323	0.02182
•		[0.0028; 0.0034]	[0.0124; 0.0482]	[0.0034; 0.0436]
σ_i	0.00465	0.0015	0.07194	0.07294
		[0.0000; 0.0028]	[0.0475; 0.0943]	[0.0560; 0.0918]
Posterior Kernel		-	-	-
Proposal approximation				
Distribution approximation		Particles	Particles	Particles
N		10,000	10,000	10,000
Mixture		l <u></u>	l	
Option		mh_replic=5,000,mh_nblck=10	$\delta = 0.9$	$\delta = 0.9$
Resampling		systematic	none	systematic

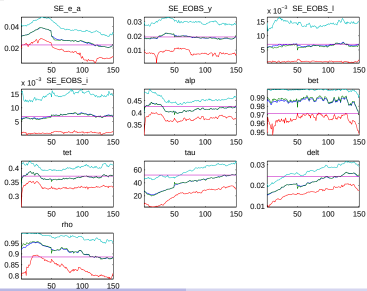
The likelihood expression
The limits of the likelihood approach
Particle MCMC

The online approach



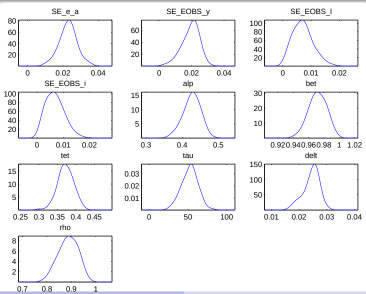
The likelihood expression The limits of the likelihood approach Particle MCMC

The online approach



The likelihood expression
The limits of the likelihood approach
Particle MCMC





In a nutshell

Filter	Proposal choice	Proposal approximation	State approximation	Resampling
Bootstrap	Transition distribution	Particles	Particles	Yes
Auxiliary	Pre-selected particles for states transition distribution	Particles	Particles	Yes,No
NL Kalman	-	-	Particles or sparse grids	-
Gaussian	Gaussian on states and errors	Sparse grid / Kalman posterior	Particles or sparse grids	Yes,No
Gaussian- Mixture	Gaussian-Mixtures on states and/or errors	Sparse grid / Kalman posterior	Particles or sparse grids	Yes,No
Conditional	Particles for states, sparse grids for errors	Kalman posterior	Particles	Yes

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