

# Graduate Macro Theory II:

## Notes on Using Dynare

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### 1 Introduction

This document will present some simple examples of how to **solve and simulate DSGE models** using Dynare. Dynare can also be used to **estimate the parameters** of DSGE models via Maximum Likelihood or Bayesian Maximum Likelihood.

Dynare is not its own program but is rather basically a collection of Matlab codes. To run Dynare, you must first install it. You can download it from the following website: <http://www.dynare.org/download>. I will only support downloads using Matlab. You need to choose a destination folder. My destination folder is in the C drive and is called “dynare”. I have version 4.4.3, the newest “stable” version (older versions will do most of the basic stuff just as well as the current version). The full path is: “C:\dynare \4.4.3 \matlab ”.

To run Dynare, you have to create .mod files (you can create these in the m-file editor, but be sure to save them as .mod, not as .m files). Then, to run your code, simply type **“dynare filename”** into the command window (or you can have this command within a separate m-file which can be in another directory so long as you call up the Dynare directory). Equivalently, you can **“set the path”** in Matlab so that you don’t always have to call up the Dynare directory.

### 2 A Neoclassical Model with Fixed Labor

Consider a simple stochastic neoclassical model with fixed labor as the laboratory. The planner’s problem can be written:

$$\begin{aligned} \max \quad & E_0 \sum_{t=0}^{\infty} \beta^t \frac{c_t^{1-\sigma} - 1}{1-\sigma} \\ \text{s.t.} \quad & \end{aligned}$$

$$k_{t+1} = a_t k_t^\alpha - c_t + (1-\delta)k_t$$

TFP is assumed to follow a mean zero AR(1) in the log:

$$\ln a_t = \rho \ln a_{t-1} + \varepsilon_t$$

The first order conditions for this problem can be characterized with three non-linear difference equations and a transversality condition:

$$\begin{aligned} c_t^{-\sigma} &= \beta E_t c_{t+1}^{-\sigma} (\alpha a_{t+1} k_{t+1}^{\alpha-1} + (1 - \delta)) \\ k_{t+1} &= a_t k_t^\alpha - c_t + (1 - \delta) k_t \\ \ln a_t &= \rho \ln a_{t-1} + \varepsilon_t \\ \lim_{t \rightarrow \infty} \beta^t c_t^{-\sigma} k_{t+1} &= 0 \end{aligned}$$

In addition, I have two auxiliary “static” variables that define output from the production function and investment from the accounting identity:

$$\begin{aligned} y_t &= a_t k_t^\alpha \\ i_t &= y_t - c_t \end{aligned}$$

I may also be interested in backing out prices (which would emerge in the equilibrium of a decentralized model). I could consider  $r_t$ ,  $R_t$ , and  $w_t$ : the real interest rate on bonds, the real rental rate on capital, and the real wage on labor (which we can still price as equaling the marginal product of labor even though labor is inelastically supplied and fixed at one). The prices are pinned down by:

$$\begin{aligned} c_t^{-\sigma} &= \beta E_t c_{t+1}^{-\sigma} (1 + r_t) \\ R_t &= \alpha a_t k_t^{\alpha-1} \\ w_t &= (1 - \alpha) a_t k_t^\alpha \end{aligned}$$

Before proceeding I must pick values of the parameters. I set  $\sigma = 1$ ,  $\alpha = 1/3$ ,  $\delta = 0.025$ ,  $\beta = 0.99$ ,  $\rho = 0.95$ , and the standard deviation of the productivity shock to 0.01.

## 2.1 A Timing Convention

There is a timing convention in Dynare that requires that we (slightly) rewrite the model. In particular, Dynare requires that predetermined variables (like the capital stock) show up as dated  $t - 1$  in the time  $t$  equations and  $t$  in the  $t + 1$  equations. This is how you “tell” Dynare if a variable is a state variable or not. As such, we need to rewrite the FOCs:

$$\begin{aligned}
c_t^{-\sigma} &= \beta E_t c_{t+1}^{-\sigma} (\alpha a_{t+1} k_t^{\alpha-1} + (1-\delta)) \\
k_t &= a_t k_{t-1}^{\alpha} - c_t + (1-\delta) k_{t-1} \\
\ln a_t &= \rho \ln a_{t-1} + \varepsilon_t \\
\lim_{t \rightarrow \infty} \beta^t c_t^{-\sigma} k_{t+1} &= 0 \\
y_t &= a_t k_{t-1}^{\alpha} \\
i_t &= y_t - c_t \\
c_t^{-\sigma} &= \beta E_t c_{t+1}^{-\sigma} (1 + r_t) \\
R_t &= \alpha a_t k_{t-1}^{\alpha-1} \\
w_t &= (1-\alpha) a_t k_{t-1}^{\alpha}
\end{aligned}$$

In other words, we essentially just lag the capital stock one period in each relevant equation.  $k_t$  is just re-interpreted as  $k_{t+1}$ .

### 3 Writing the Dynare Code

With the first order conditions in hand, we are now ready to begin to write our dynare code (which occurs in the .mod file). The dynare code will not run unless each entry is followed by a semicolon to suppress the output.

The first step is to **declare the variables of the model**. Dynare automatically sorts variables into the following order: static variables (i.e. only enter FOCs at time  $t$ ), purely predetermined variables (only enter FOCs at  $t$  and  $t-1$ ), variables that are both predetermined and forward-looking (appear at  $t-1$ ,  $t$ , and  $t+1$ ), and variables that are purely forward-looking (only appear at dates  $t$  and  $t+1$ ). In our model, output is static, capital is purely predetermined, technology is both predetermined and forward-looking (since future TFP shows up in the Euler equation), and consumption is forward-looking. It makes sense to declare the variables in this order. The first non-comment command in your Dynare code should be: “var” followed by the endogenous variable names and culminating with a semicolon. For the purposes of naming things, everything other than the shock(s) is considered to be endogenous. My first line of code looks like:

```
1 var y I k a c w R r;
```

The next line of code specifies the exogenous variables. These are just the shocks, and in this model I only have one:

```
1 varexo e;
```

The next step is to declare the parameters of the model. You simply type “parameters” followed by parameter names:

```
1 parameters alpha beta Δ rho sigma sigmae;
```

Immediately following this command, you need to **specify numerical values for these parameters:**

```
1 alpha = 1/3;
2 beta = 0.99;
3 Δ = 0.025;
4 rho = 0.95;
5 sigma = 1;
6 sigmae = 0.01;
```

Now that you’ve named variables and specified parameter values, it’s now time to **“declare the model”**. To do this, you type in “model;” followed by the **first order conditions, constraints, identities**, etc., followed by “end;”. We typically want to approximate the solutions of the natural logs of the variables, so that impulse responses, etc. are all in percentage terms. Dynare will do approximations of the levels of the variables. To get it to **do linear approximation of the logs of the variables, you need to specify the variables as “exp( $x$ )”**. This way the variable  $x$  is interpreted as the log of the variable of interest, while  $\exp(x)$  is the level (since the exponential and log are inverse functions), which is what shows up in most of the FOCs. Then you just type in the first order conditions. If a variable appears dated  $t$ , then you just type  $x$ . If a variable is  $t + 1$ , you type  $x(+1)$ ; if it’s  $t - 1$ , then  $x(-1)$ . If, for whatever reason, you need variables beyond one period away to appear, you could define auxiliary state variables to make the system work out this way. My first order conditions for the above model look like:

```
1 model;
2
3 exp(c)^(-sigma) = beta*exp(c(+1))^(-sigma)*(alpha*exp(a(+1))*exp(k)^(alpha-1) + (1-Δ));
4 exp(y) = exp(a)*exp(k(-1))^(alpha);
5 exp(k) = exp(a)*exp(k(-1))^(alpha) - exp(c) + (1-Δ)*exp(k(-1));
6 a = rho*a(-1) + e;
7 exp(y) = exp(c) + exp(I);
8 exp(c)^(-sigma) = beta*exp(c(+1))^(-sigma)*(1+r);
9 exp(R) = alpha*exp(a)*exp(k(-1))^(alpha-1);
10 exp(w) = (1-alpha)*exp(a)*exp(k(-1))^(alpha);
11
12
13 end;
```

Dynare interprets all of the **underlying variables here as logs**. Since I specified the stochastic process for  $a_t$  in the log, I do not need to write it using the exponential. Dynare will solve for the

steady **state numerically for you.** Alternatively, you can create **a steady state file to solve for the steady state** “by hand” (more on this later). To get it to do this you need to give it guess of the steady state. If you give it bad guesses, Dynare may not converge and you can get an error. When giving initial values, remember that Dynare is interpreting all the variables as logs. Hence, if you know that steady state capital is 30, you want to give an initial value of capital somewhere near  $\ln 30$ , not 30. you begin this part of the code with the command “initval;”, followed by the initial values/guesses of all the endogenous variables, capped off with “end”. My code for this part looks like:

```
1 initval;
2 k = log(30);
3 y = log(3);
4 c = log(2.5);
5 I = log(0.5);
6 a = 0;
7 r = (1/beta) - 1;
8 R = log((1/beta) - (1-Δ));
9 w = log(1);
10 end;
```

The next step is to specify the variance of the shocks. This part of the code starts with “shocks;”, followed by a specification of the *variance* (not standard deviation), followed by “end;”:

```
1 var e = sigmae^2;
```

In the next step you simply type in ~~“steady;”. This command calculates the steady state values~~ of the endogenous variables of the model:

```
1 steady;
```

The next command is the payoff. It’s the “stoch\_simul” command, which is what solves the model, produces the policy functions, and generates impulse responses functions and unconditional second moments. There are a number of options following this command. If you just type in “stoch\_simul;”, it’s going to give you the default output. **The default output** is: (1) steady state values of endogenous variables; (2) a model summary, which counts variables by type; (3) covariance matrix of shocks (which in the example I’m giving is a scalar); (4) the policy and transition functions (in state space notation); (5) theoretical first and second moments; (6) a theoretical correlation matrix; (7) theoretical autocovariances up to order 5; and (8) impulse responses. By default, Dynare does a second order approximation about the steady state; the second order approximation involves “cross terms” , and so the policy/transition functions look more complicated than you’re used to seeing.

There are a number of options one can include behind “stoch\_simul;” (to do this you type

`“stoch_simul(options);”` . There are several of these that you can read about in the manual, but the more important ones for our purposes are:

- `hp_filter = integer`: this will produce theoretical moments (variances, covariances, autocorrelations) after HP filtering the data (the default is to apply no filter to the data). We typically want to look at HP filtered moments, so this is an option you’ll want to use. The integer is a number (i.e. 1600) corresponding to the penalty parameter in the HP filter. So, for example, typing `“stoch_simul(hp_filter=1600);”` will produce theoretical (i.e. analytical) moments of the HP filtered data. The only problem here is that it will not simultaneously do HP filtering of simulated data; you can either get moments from simulated data or analytical HP filtered moments
- `irf = integer`: this will change the number of periods plotted in the impulse response functions. The default value is 40. To suppress printing impulse responses altogether, type in 0 for the number of horizons. For example, to see impulse responses for only 20 periods, type `“stoch_simul(irf=20);”`.
- `nocorr: this` will mean it will not print a matrix of correlations in the command window
- `nofunctions`: this will suppress printing the state space coefficients of the solution in the command window
- `nomoments: this` will suppress printing of moments
- `noprint: this` will suppress printing of any output at all
- `order = 1, 2, or 3`: this tells Dynare the order of the (log) approximation. The default is a second order approximation. Hence, typing `“order=1”` will have it do a linear approximation.
- `periods = integer`: Dynare’s default is to produce analytical/theoretical moments of the variables. Having periods not equal to zero will instead have it simulate data and take the moments from the simulated data. By default, Dynare drops the first 100 values from a simulation, so you need to give it a number of periods greater than 100 for this to work. Hence, typing `“stoch_simul(periods=300);”` will produce moments based on a simulation with 200 variables.
- `drop = integer`: You can change the number of observations to drop from the simulations. For example, typing `“stoch_simul(drop=0);”` will result in no observations being dropped in the simulations.
- `simul_seed = integer`: sets the seed used in the random number generator for the simulations.

For example, typing `“stoch_simul(nofunctions, hp_filter=1600, order=1, irf=20);”` will suppress the policy function output, will produce analytical HP filtered moments, will do a first order approximation, and will plot 20 periods in the impulse responses.

Dynare will produce **policy and transition functions** for the endogenous variables of the model in state space form. Letting  $s_t$  be a  $m \times 1$  vector of states (here  $m = 2$ ) and  $x_t$  be a  $n \times 1$  vector of controls (here I include “static” controls like output as well as dynamic controls; you don’t need to eliminate the static controls to use Dynare, because Dynare can figure out that they’re static/redundant), which in this model is  $n = 6$ . The state space representation of the system can be written:

$$\begin{aligned}s_t &= As_{t-1} + B\varepsilon_t \\ x_t &= \Phi s_t\end{aligned}$$

Above  $A$  is  $m \times m$ , and can be found as discussed previously in class.  $\varepsilon_t$  is  $w \times 1$ ; so in this model  $w = 1$ ; and  $B$  is  $m \times w$ .  $\Phi$  is the policy function, and is  $n \times m$ . Dynare writes the system out by substituting the state transition equation into the policy function as follows:

$$x_t = \Phi As_{t-1} + \Phi B\varepsilon_t$$

It is easily verified that these matrix multiplications are permissible;  $\Phi$  is  $n \times m$  and  $A$  is  $m \times m$ . Hence  $\Phi A$ , call it  $C$ , is  $n \times m$ .  $B$  is  $m \times w$ ; hence  $\Phi B$  is  $n \times w$ . Call  $\Phi B = D$ . Then we can write **out the full system as**:

$$\begin{aligned}s_t &= As_{t-1} + B\varepsilon_t \\ x_t &= Cs_{t-1} + D\varepsilon_t\end{aligned}$$

We can combine this into one expression by denoting  $Y_t = [s_t \ x_t]'$ ,  $\Psi = [A \ C]'$ , and  $\Omega = [B \ D]'$ :

$$Y_t = \Psi s_{t-1} + \Omega \varepsilon_t$$

### 3.1 The Full .mod File

Below is my code in the .mod file in its entirety. Note that you can comment in a .mod file just like you can in a regular .m file.

```
1 var y I k a c w R r;
2
3 varexo e;
4
5 parameters alpha beta Δ rho sigma sigmae;
```

```

6
7  alpha = 1/3;
8  beta = 0.99;
9  Δ = 0.025;
10 rho = 0.95;
11 sigma = 1;
12 sigmae = 0.01;
13
14 model;
15
16 exp(c)^(-sigma) = beta*exp(c(+1))^( -sigma)*(alpha*exp(a(+1))*exp(k)^(alpha-1) + (1-Δ));
17 exp(y) = exp(a)*exp(k(-1))^(alpha);
18 exp(k) = exp(a)*exp(k(-1))^(alpha) - exp(c) + (1-Δ)*exp(k(-1));
19 a = rho*a(-1) + e;
20 exp(y) = exp(c) + exp(I);
21 exp(c)^(-sigma) = beta*exp(c(+1))^( -sigma)*(1+r);
22 exp(R) = alpha*exp(a)*exp(k(-1))^(alpha-1);
23 exp(w) = (1-alpha)*exp(a)*exp(k(-1))^(alpha);
24
25 end;
26
27 initval;
28 k = log(30);
29 y = log(3);
30 c = log(2.5);
31 I = log(0.5);
32 a = 0;
33 r = (1/beta) - 1;
34 R = log((1/beta) - (1-Δ));
35 w = log(1);
36 end;
37
38 shocks;
39 var e = sigmae^2;
40 end;
41
42 steady;
43
44 stoch_simul(order=1, irf=20);

```

## 4 Running Dynare

To run Dynare you can't hit F5 or type the name of the .mod file into the command prompt. You have to type **"dynare filename"** (without the .mod on the end of the file name). For this to run you must either be in the same directory where your Dynare files are stored, you must have set the path of where your Dynare files are stored, or you must type "addpath (directory)" where the directory is the location of your Dynare files. It is also helpful to add "noclearall" to the declaration



so that you are typing in “dynare filename noclearall”. Otherwise Dynare will automatically erase all previous output, which you may not want. Another option I often include is “nolog” which tells Dynare to not produce a “log” file. I have found in the past that the creation of log files can cause crashes when you’re doing loops.

So as to do other things, I usually write a .m file to go along with my .mod file. My code for the .m file is simple:

```

1 clear all;
2 close all;
3
4 % execute a Dynare file
5
6 % specify parameters
7
8 beta = 0.99;
9 alpha = 1/3;
10 sigma = 1;
11 sigmae = 0.01;
12 rho = 0.9;
13 Δ = 0.025;
14
15 save param.nc alpha beta Δ rho sigma sigmae
16
17 dynare basic.nc_dynare.alt noclearall nolog

```

Here I’m saving the parameters as a .mat file. Dynare allows you to do this, and you can modify the .mod file (as below) to accommodate this. This is particularly useful if you want to produce moments for different parameter configurations that you might want to loop over.

```

1 var y I k a c w R r;
2
3 varexo e;
4
5 parameters alpha beta Δ rho sigma sigmae;
6
7 load param.nc;
8 set_param_value('alpha',alpha);
9 set_param_value('beta',beta);
10 set_param_value('sigma',sigma);
11 set_param_value('Δ',Δ);
12 set_param_value('rho',rho);
13 set_param_value('sigmae',sigmae);
14
15 model;
16
17 exp(c)^(-sigma) = beta*exp(c(+1))^(sigma)*(alpha*exp(a(+1))*exp(k)^(alpha-1) + (1-Δ));

```

```

18 exp(y) = exp(a)*exp(k(-1))^(alpha);
19 exp(k) = exp(a)*exp(k(-1))^(alpha) - exp(c) + (1-Δ)*exp(k(-1));
20 a = rho*a(-1) + e;
21 exp(y) = exp(c) + exp(I);
22 exp(c)^(-sigma) = beta*exp(c(+1))^(sigma)*(1+r);
23 exp(R) = alpha*exp(a)*exp(k(-1))^(alpha-1);
24 exp(w) = (1-alpha)*exp(a)*exp(k(-1))^(alpha);
25
26 end;
27
28 initval;
29 k = log(30);
30 y = log(3);
31 c = log(2.5);
32 I = log(0.5);
33 a = 0;
34 r = (1/beta) - 1;
35 R = log((1/beta) - (1-Δ));
36 w = log(1);
37 end;
38
39 shocks;
40 var e = sigmae^2;
41 end;
42
43 steady;
44
45 stoch_simul(order=1, irf=20);

```

You can see how I modified the code in the “parameter” block to accommodate the loading in of saved parameters.

By default, Dynare computes the steady state numerically – you give it initial values (if you don’t give it initial values it uses values by default, which I think are 0 or 1), and it numerically computes the steady state. This is fine, but can be prone to problems. First, if you give it bad initial values it may not find the steady state, at which point your program crashes. Second, you may encounter situations in which you want to solve the model for different parameter values which could impact the steady state. What may be **good initial values** for one set of parameters may be bad initial values for another.

There is an easy way to get around this issue – write your own steady state file which solves the steady state analytically for any values of the parameters. You simply name the file the same thing as your .mod file, appended with “\_steadystate.m” (note the steady state file is an .m file, not a .mod file). The **structure “M” that** Dynare produces saves the parameter values; in your steady state file you read the parameters from “M,” solve for the steady state values of the variables of the model, and then create matrix called “ys” which has the steady state values. My steady state code is given below:

```

1 function [ys,check] = basic_nc_dynare_ss_steadystate(ys,exe);
2
3 global M_
4
5 alpha = M_.params(1);
6 beta = M_.params(2);
7 Δ = M_.params(3);
8 rho = M_.params(4);
9 sigma = M_.params(5);
10 sigmae = M_.params(6);
11
12 k = (alpha/(1/beta - (1-Δ)))^(1/(1-alpha));
13 y = k^(alpha);
14 I = Δ*k;
15 c = y - I;
16 a = 1;
17 w = (1-alpha)*k^(alpha);
18 R = alpha*k^(alpha-1);
19 r = (1/beta) - 1;
20
21 check = 0;
22
23 ys = [log(y);
24       log(I);
25       log(k);
26       log(a);
27       log(c);
28       log(w);
29       log(R);
30       r];

```

You have to follow this structure (e.g. for whatever reason you need the “check = 0” line in there). But once you have this, you can eliminate the “initval” block of your .mod file and it’ll automatically recognize the steady state file.

Finally, something else that comes in handy is to note that Dynare produces *a lot* of output when you run it. One file it produces is a .m file with the same name as your .mod file. Once you’ve run the .mod file once, you can then just type in the name of the .m file into the command window in Matlab and it’ll produce all the same output. This turns out to be substantially faster than typing “dynare file name” because there is all this initialization that Dynare does that slows it down (you will see this immediately below).

## 5 Dynare Output

Dynare will produce a bunch of output in the command window for you (unless you include options discussed above to suppress it). Below is the output that I get:

```
Configuring Dynare ...
[mex] Generalized QZ.
[mex] Sylvester equation solution.
[mex] Kronecker products.
[mex] Sparse kronecker products.
[mex] Local state space iteration (second order).
[mex] Bytecode evaluation.
[mex] k-order perturbation solver.
[mex] k-order solution simulation.
[mex] Quasi Monte-Carlo sequence (Sobol).
[mex] Markov Switching SBVAR.
```

```
Starting Dynare (version 4.4.2).
Starting preprocessing of the model file ...
Found 8 equation(s).
Evaluating expressions...done
Computing static model derivatives:
- order 1
Computing dynamic model derivatives:
- order 1
Processing outputs ...done
Preprocessing completed.
Starting MATLAB/Octave computing.
```

#### **STEADY-STATE RESULTS:**

```
y    1.12546
I    -0.312508
k    3.37637
a     0
c    0.854424
w    0.719993
R    -3.34953
r    0.010101
```

#### **MODEL SUMMARY**

```
Number of variables:      8
Number of stochastic shocks: 1
```

Number of state variables: 2  
Number of jumpers: 2  
Number of static variables: 5

#### MATRIX OF COVARIANCE OF EXOGENOUS SHOCKS

Variables e  
e 0.000100

#### POLICY AND TRANSITION FUNCTIONS

	y	I	k	a	
Constant	1.125457	-0.312508	3.376372	0	0.8
k(-1)	0.333333	-0.502551	0.962436	0	0.8
a(-1)	0.900000	3.139964	0.078499	0.900000	0.2
e	1.000000	3.488849	0.087221	1.000000	0.2

#### THEORETICAL MOMENTS

VARIABLE	MEAN	STD. DEV.	VARIANCE
y	1.1255	0.0286	0.0008
I	-0.3125	0.0743	0.0055
k	3.3764	0.0275	0.0008
a	0.0000	0.0229	0.0005
c	0.8544	0.0194	0.0004
w	0.7200	0.0286	0.0008
R	-3.3495	0.0212	0.0005
r	0.0101	0.0007	0.0000

#### MATRIX OF CORRELATIONS

Variables	y	I	k	a	c	w	R	r
y	1.0000	0.9017	0.7567	0.9600	0.8569	1.0000	0.4202	0.3167
I	0.9017	1.0000	0.3997	0.9867	0.5499	0.9017	0.7712	0.6957
k	0.7567	0.3997	1.0000	0.5433	0.9854	0.7567	-0.2752	-0.3804
a	0.9600	0.9867	0.5433	1.0000	0.6783	0.9600	0.6576	0.5697
c	0.8569	0.5499	0.9854	0.6783	1.0000	0.8569	-0.1076	-0.2175

w	1.0000	0.9017	0.7567	0.9600	0.8569	1.0000	0.4202	0.3167
R	0.4202	0.7712	-0.2752	0.6576	-0.1076	0.4202	1.0000	0.9938
r	0.3167	0.6957	-0.3804	0.5697	-0.2175	0.3167	0.9938	1.0000

#### COEFFICIENTS OF AUTOCORRELATION

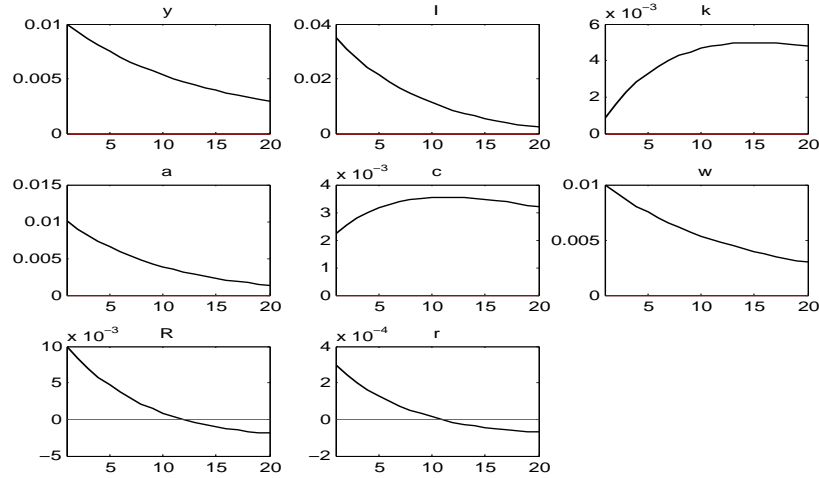
Order	1	2	3	4	5
y	0.9367	0.8784	0.8246	0.7748	0.7289
I	0.8827	0.7777	0.6839	0.6001	0.5252
k	0.9980	0.9925	0.9840	0.9730	0.9597
a	0.9000	0.8100	0.7290	0.6561	0.5905
c	0.9927	0.9826	0.9702	0.9559	0.9398
w	0.9367	0.8784	0.8246	0.7748	0.7289
R	0.8768	0.7668	0.6686	0.5811	0.5031
r	0.8868	0.7853	0.6945	0.6133	0.5406

Total computing time : 0h00m01s

Several comments on the nature of the output. First, Dynare starts with a “model summary.” This includes variables by “type” but note that the types (what Dynare calls states, jumpers, and static) do not necessarily sum to the total number of variables. Dynare counts things which show up in the equilibrium conditions just dated  $t$  as static variables; things which show up with a  $t - 1$  as states, and things which show up with a  $t + 1$  as jumpers. Something could show up as both a state and a jumper – here,  $a$  appears lagged (as a state) and with a “+1” in the Euler equation (jumper).

The next part of the Dynare output are the policy and transition functions. The “constant” in the policy and transition functions output is just the steady state values; the coefficients of the (modified) state space representation are easy to understand. In addition, Dynare produces means (steady state values), volatilities (standard deviations), and variances for each endogenous variable. It also produces the matrix of correlations as well as coefficients of autocorrelation at different lags. If there are multiple shocks, it will also produce a variance decomposition telling you how important each shock is – using the terminology I’ve used, this is an “unconditional” variance decomposition in the sense of showing the fraction of the total variance of each variable. Given the options I specified, all of these moments are theoretical (in the sense of being analytical, not based on a simulation).

Dynare also produces impulse responses to the shocks. These are shown here:



There is one interpretive issue in the IRFs of which you need to be aware. Recall that Dynare interprets  $k_t$  as  $k_{t+1}$ . Hence, the impulse response of  $k_t$  that it produces is really the impulse response of  $k_{t+1}$ . To get the impulse responses of  $k_t$ , you need to (i) add a zero as the impact response (remember the capital stock can't react on impact as it is predetermined) and (ii) set the response of  $k_t$  at horizon  $h$  equal to the response of what Dynare gives as the impulse response in period  $h + 1$ .

## 6 Where is the Output Stored?

Dynare is great at producing lots of different output right there for you to see. It's a little more challenging to find where your output is stored.

Dynare will store IRFs for you under the following nomenclature: **"x\_e"** denotes the impulse response of variable  $x$  to shock  $e$ . The coefficients of the state space representation are a bit harder to find. In terms of the notation introduced above, the coefficients of the  $\Psi$  matrix are stored in **"oo\_dr.ghx"**; the coefficients of the  $\Omega$  matrix are stored in **"oo\_dr.ghu"**; and the steady state values are stored in **"oo\_dr.js"**. One irritation with Dynare is that it does not store coefficients of the state space representation in the same order you declare variables (whereas the output produced in the command window is based on what is called the "order of declaration" or the order in which you declare the variables). The **"DR" ordering (for "decision rule")** is based on the following ordering of variables by type: static variables (only dated  $t$ ), backward-looking variables (dated  $t$  and  $t - 1$ ), mixed variables (dated  $t - 1$ ,  $t$ , and  $t + 1$ ), and forward-looking variables (variables dated  $t$  and  $t + 1$ ). The stored output **"oo\_dr.inv.order\_var"** contains the the inverse mapping – so if a variable is declared 2nd, **"oo\_dr.inv.order\_var(2)"** will give you the DR order for that variable.

If you want to use the coefficients of the state space representation to do your own simulations, etc., it is helpful to first declare variable orderings in your own code according to the order in which you declared them:

```

1 p_y = 1;
2 p_I = 2;
3 p_k = 3;
4 p_a = 4;
5 p_c = 5;
6 p_w = 6;
7 p_R = 7;
8 p_r = 8;

```

Then I can create matrixes for the state space representation as follows:

```

1 % state space representation: S(t) = A*S(t-1) + B*e(t), X(t) = C*S(t-1) +
2 % D*e(t);
3
4 A = [oo.dr.ghx(oo.dr.inv_order_var(p_k),:);
5      oo.dr.ghx(oo.dr.inv_order_var(p_a),:)]';
6
7 B = [oo.dr.ghu(oo.dr.inv_order_var(p_k),:);
8      oo.dr.ghu(oo.dr.inv_order_var(p_a),:)]';
9
10 C = [oo.dr.ghx(oo.dr.inv_order_var(p_y),:);
11      oo.dr.ghx(oo.dr.inv_order_var(p_I),:);
12      oo.dr.ghx(oo.dr.inv_order_var(p_c),:);
13      oo.dr.ghx(oo.dr.inv_order_var(p_w),:);
14      oo.dr.ghx(oo.dr.inv_order_var(p_R),:);
15      oo.dr.ghx(oo.dr.inv_order_var(p_r),:)]';
16
17 D = [oo.dr.ghu(oo.dr.inv_order_var(p_y),:);
18      oo.dr.ghu(oo.dr.inv_order_var(p_I),:);
19      oo.dr.ghu(oo.dr.inv_order_var(p_c),:);
20      oo.dr.ghu(oo.dr.inv_order_var(p_w),:);
21      oo.dr.ghu(oo.dr.inv_order_var(p_R),:);
22      oo.dr.ghu(oo.dr.inv_order_var(p_r),:)]';

```

Then with this, I can create impulse responses on my own, and can verify that they check out with what Dynare produces already:

```

1 % compute impulse responses by hand
2 H = 20;
3 Sirf = zeros(2,H);
4 Xirf = zeros(6,H);
5
6 Sirf(:,1) = B*sigmae;
7 Xirf(:,1) = D*sigmae;
8
9 for j = 2:H
10     Sirf(:,j) = A*Sirf(:,j-1);

```



```

11      Xirf(:,j) = C*Sirf(:,j-1);
12  end

```

It is straightforward to verify that these impulse responses are the same as what Dynare produces.

It is helpful to know where these are in the event that you want to conduct your own simulation. I can simulate out variables just as before by drawing shocks from some distribution. For the simulation I assume that everything begins in steady state, so that the initial conditions are all 0 (recall that the variables are deviations about steady state). Here is some code to do that:

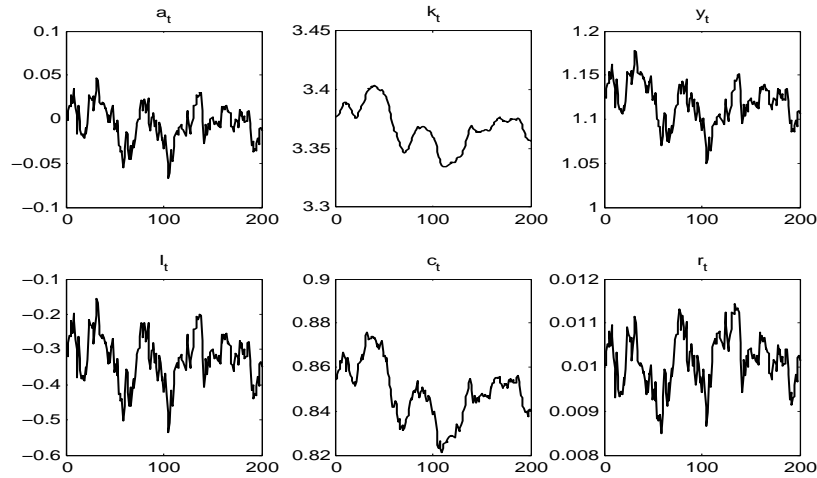
```

1  % compute a simulation.  First draw shocks
2  randn('seed',666)
3  T = 200; % number of periods to simulate
4  e = sigmae*randn(1,T);
5
6  Ssim = zeros(2,T);
7  Xsim = zeros(6,T);
8
9  % assume initial state is SS
10 Ssim(:,1) = B*e(1,1);
11 for j = 2:T
12     Ssim(:,j) = A*Ssim(:,j-1) + B*e(1,j);
13     Xsim(:,j) = C*Ssim(:,j-1) + D*e(1,j);
14 end

```

As written, the interpretation of the variables in the two matrixes are as deviations about steady state, so these ought to be mean-zero (in a large enough sample). One might want to add back in the means, which in a linear approximation is just the steady state values. The steady state values are stored in “oo\_.dr.ys” – but the steady state values are stored in the order you declared the variables, which is different than the stored coefficients of the state space representation.

Here is a figure of the simulated time paths of these variables:



The simulations have some of the features we might expect – consumption is “smoother” than output, which is less volatile than investment. The capital stock evolves very smoothly as well.

Dynare will also directly do simulations for you if you want. For example, if I type in “`stoch_simul(order=1,irf=40,periods=200)`” it will generate a simulation of 200 periods of the variables of the model. If you use this command then it produces moments (standard deviations, correlations, and autocorrelations) off of the simulated data, as opposed to producing “theoretical” moments using the coefficients of the state space representation **(theoretical in the sense that you can calculate analytic moments of the state-space representation because it is linear).**

Dynare will also store the second moments statistics that it produces. “`oo.mean`” will produce the means of the variables; “`oo.var`” produces the variance-covariance matrix (the standard deviations are the square roots of the diagonal elements). “`oo.autocorr`” will produce autocorrelations – “`oo.autocorr{i}(k,l)`” will produce the  $i$ th autocorrelation coefficient between the variables  $k$  and  $l$  – e.g.  $corr(y_t^l, y_{t-i}^l)$ . So the first order autocorrelation coefficient of variable  $j$  would be “`oo.autocorr{1}(j,j)`.” The field “`oo.gamma_y{AR+2}`” will give you the variance decomposition (only if there is more than one shock), where AR is the number of AR lags it uses (the default is AR = 5).

## 7 Higher Order Approximations

Dynare will use **higher order perturbations (based on 2nd or 3rd order approximations).** In fact, by default Dynare solves models using a 2nd order approximation (you have to change the options in “`stoch_simul`” to get a first order approximation. Higher order approximations have several potential advantages. First, higher order approximations are more appropriate for welfare analysis. Second, higher order approximations are going to be more accurate. Third, higher order approximations can allow you to examine state-dependence (e.g. do impulse responses depend on the initial state in which a shock hits). Fourth, higher order approximations are necessary if you want

to think about “second moment” shocks (e.g. shocks to the variance of a shock . . . you actually need a third order approximation for this to introduce interesting dynamics).

For a second order approximation, Dynare will produce the following output for the **state space** representation:

$$s_t = \frac{1}{2}\Upsilon_{s,0} + As_{t-1} + B\epsilon_t + \frac{1}{2}\Upsilon_{s,1}(s_{t-1} \otimes s_{t-1}) + \frac{1}{2}\Upsilon_{s,2}(\epsilon_t \otimes \epsilon_t) + \Upsilon_{s,3}(s_{t-1} \otimes \epsilon_t) \quad (1)$$

$$x_t = \frac{1}{2}\Upsilon_{x,0} + Cs_{t-1} + D\epsilon_t + \frac{1}{2}\Upsilon_{x,1}(s_{t-1} \otimes s_{t-1}) + \frac{1}{2}\Upsilon_{x,2}(\epsilon_t \otimes \epsilon_t) + \Upsilon_{x,3}(s_{t-1} \otimes \epsilon_t) \quad (2)$$

The new coefficients are the higher order terms; I call them  $\Upsilon$  and index them by  $s$  or  $x$  (for state or non-state). It is *not* a typo that a  $\frac{1}{2}$  term doesn’t appear in the last term (think about taking a second order approximation . . . the cross terms are symmetric and show up twice). A first order approximation is a special case of this where all the  $\Upsilon$  coefficient matrixes are zero. Note also that the first order coefficients,  $A$ ,  $B$ ,  $C$ , and  $D$  are the same whether the second order terms are zero or not. The operator  $\otimes$  is the Kronecker product. I will not go into the details of how these coefficients are found.

The Dynare output for the second order approximation is given below:

#### STEADY-STATE RESULTS:

```
y    1.12546
I    -0.312508
k    3.37637
a    0
c    0.854424
w    0.719993
R    -3.34953
r    0.010101
```

#### MODEL SUMMARY

```
Number of variables:      8
Number of stochastic shocks: 1
Number of state variables: 2
Number of jumpers:       2
Number of static variables: 5
```

#### MATRIX OF COVARIANCE OF EXOGENOUS SHOCKS

Variables	e
e	0.000100

#### POLICY AND TRANSITION FUNCTIONS

		y	I	k	a	
Constant	1.125457	-0.312578	3.376370	0	0.8	0.8
(correction)	0	-0.000070	-0.000002	0	0.0	0.0
k(-1)	0.333333	-0.502551	0.962436	0	0.5	0.5
a(-1)	0.900000	3.139964	0.078499	0.900000	0.2	0.2
e	1.000000	3.488849	0.087221	1.000000	0.2	0.2
k(-1),k(-1)	0	-0.536347	0.014107	0	0.0	0.0
a(-1),k(-1)	0	2.684454	-0.047889	0	-0.0	-0.0
a(-1),a(-1)	0	-3.408636	0.034945	0	0.0	0.0
e,e	0	-4.208192	0.043142	0	0.0	0.0
k(-1),e	0	2.982727	-0.053210	0	-0.0	-0.0
a(-1),e	0	-7.574746	0.077656	0	0.0	0.0

#### APPROXIMATED THEORETICAL MOMENTS

VARIABLE	MEAN	STD. DEV.	VARIANCE
y	1.1256	0.0286	0.0008
I	-0.3145	0.0743	0.0055
k	3.3768	0.0275	0.0008
a	0.0000	0.0229	0.0005
c	0.8547	0.0194	0.0004
w	0.7201	0.0286	0.0008
R	-3.3498	0.0212	0.0005
r	0.0101	0.0007	0.0000

#### APPROXIMATED MATRIX OF CORRELATIONS

Variables	y	I	k	a	c	w	R	r
y	1.0000	0.9017	0.7567	0.9600	0.8569	1.0000	0.4202	0.3167
I	0.9017	1.0000	0.3997	0.9867	0.5499	0.9017	0.7712	0.6957
k	0.7567	0.3997	1.0000	0.5433	0.9854	0.7567	-0.2752	-0.3804
a	0.9600	0.9867	0.5433	1.0000	0.6783	0.9600	0.6576	0.5697

c	0.8569	0.5499	0.9854	0.6783	1.0000	0.8569	-0.1076	-0.2175
w	1.0000	0.9017	0.7567	0.9600	0.8569	1.0000	0.4202	0.3167
R	0.4202	0.7712	-0.2752	0.6576	-0.1076	0.4202	1.0000	0.9938
r	0.3167	0.6957	-0.3804	0.5697	-0.2175	0.3167	0.9938	1.0000

#### APPROXIMATED COEFFICIENTS OF AUTOCORRELATION

Order	1	2	3	4	5
y	0.9367	0.8784	0.8246	0.7748	0.7289
I	0.8827	0.7777	0.6839	0.6001	0.5252
k	0.9980	0.9925	0.9840	0.9730	0.9597
a	0.9000	0.8100	0.7290	0.6561	0.5905
c	0.9927	0.9826	0.9702	0.9559	0.9398
w	0.9367	0.8784	0.8246	0.7748	0.7289
R	0.8768	0.7668	0.6686	0.5811	0.5031
r	0.8868	0.7853	0.6945	0.6133	0.5406

Total computing time : 0h00m02s

What Dynare labels “correction” is the coefficient  $\Upsilon_{s,0}$  and  $\Upsilon_{x,0}$ . The coefficients on  $k(-1)$ ,  $a(-1)$  and  $e$  are the values in the  $A$ ,  $B$ ,  $C$ , and  $D$  matrixes we had above. The coefficients on  $k(-1)$ ,  $k(-1)$ ,  $a(-1)$ ,  $k(-1)$ , and  $a(-1)$ ,  $a(-1)$  are the elements  $\Upsilon_{s,1}$  and  $\Upsilon_{x,1}$ . The coefficients on  $e$ ,  $e$  are the  $\Upsilon_{s,2}$  and  $\Upsilon_{x,2}$  matrixes, while the  $k(-1)$ ,  $e$  and  $a(-1)$ ,  $e$  coefficients are the elements of  $\Upsilon_{s,3}$  and  $\Upsilon_{x,3}$ . The rest of the output for Dynare is the same as before, although these are “approximate theoretical moments” sense the model is non-linear. Note that what Dynare produces in the command window already includes the multiplication by  $\frac{1}{2}$ : what stores for you does not have this, so if you want to do your own simulations, you need to take this into account.

Dynare will produce impulse responses just as before. How does it do this in a non-linear model? It does so by simulation. In a nutshell, it will produce two simulations (using the same draw of shocks), but in one of the periods will add one standard deviation to the shock (or one standard deviation to one of the shocks in a multi-shock model). This is somewhat tricky because the impulse responses here will depend upon the initial value of the state. S

Here is what Dynare does. It will produce  $T$  different simulations with  $H_1 + H_2$  periods in each simulation, starting from the steady state. It will then average over the  $T$  simulations. Then it will repeat this exact same process (using the same draw of shocks), but in period  $H_1 + 1$  it will add one standard deviation to the shock (or add one standard deviation to one of the shocks in a one shock model). Then you average across  $T$  in the second simulation. Then for periods  $H_1 + 1$  through  $H_1 + H_2$ , you subtract the difference between the second and first simulations, and this will give you the impulse response function. There are two components to this process. The first

$H_1$  periods in essence averages over the initial state vector – if you knew the state from which you wanted to start the impulse response function, you could set  $H_1 = 0$ . In this sense what Dynare produces are in a sense “average” impulse response. The second component is the typical approach for simulating an impulse response function in a non-linear model.

Dynare stores in the coefficients in a similar manner as in a first order approximation. I can retrieve the  $A$ ,  $B$ ,  $C$ , and  $D$  matrixes as before. The  $\Upsilon_{s,1}$  and  $\Upsilon_{x,1}$  coefficients are stored in “oo.dr.ghxx,” the  $\Upsilon_{s,2}$  and  $\Upsilon_{x,2}$  coefficients in “oo.dr.ghuu,” the  $\Upsilon_{s,3}$  and  $\Upsilon_{x,3}$  coefficients in “oo.dr.ghxu,” and the  $\Upsilon_{s,0}$  and  $\Upsilon_{x,0}$  coefficients in “oo.dr.ghs2.” I retrieve the coefficients using the code below:

```

1 [tn,sn] = size(oo.dr.ghx); % dimensions, tn is total number of variables,
2 % sn is number of states
3 A = zeros(sn,sn);
4 B = zeros(sn,1);
5 Us0 = zeros(sn,1);
6 Us1 = zeros(sn,sn^2);
7 Us2 = zeros(sn,1);
8 Us3 = zeros(sn,sn*1); % 1 is the number of shocks
9
10 C = zeros(tn-sn,sn);
11 D = zeros(tn-sn,1);
12 Us0 = zeros(tn-sn,1);
13 Us1 = zeros(tn-sn,sn^2);
14 Us2 = zeros(tn-sn,1);
15 Us3 = zeros(tn-sn,sn*1); % 1 is the number of shocks
16
17 A = [oo.dr.ghx(oo.dr.inv_order_var(p_k),:);
18      oo.dr.ghx(oo.dr.inv_order_var(p_a),:)]';
19
20 B = [oo.dr.ghu(oo.dr.inv_order_var(p_k),:);
21      oo.dr.ghu(oo.dr.inv_order_var(p_a),:)]';
22
23 C = [oo.dr.ghx(oo.dr.inv_order_var(p_y),:);
24      oo.dr.ghx(oo.dr.inv_order_var(p_I),:);
25      oo.dr.ghx(oo.dr.inv_order_var(p_c),:);
26      oo.dr.ghx(oo.dr.inv_order_var(p_w),:);
27      oo.dr.ghx(oo.dr.inv_order_var(p_R),:);
28      oo.dr.ghx(oo.dr.inv_order_var(p_r),:)]';
29
30 D = [oo.dr.ghu(oo.dr.inv_order_var(p_y),:);
31      oo.dr.ghu(oo.dr.inv_order_var(p_I),:);
32      oo.dr.ghu(oo.dr.inv_order_var(p_c),:);
33      oo.dr.ghu(oo.dr.inv_order_var(p_w),:);
34      oo.dr.ghu(oo.dr.inv_order_var(p_R),:);
35      oo.dr.ghu(oo.dr.inv_order_var(p_r),:)]';
36

```

```

37 Us1 = [oo.dr.ghxx(oo.dr.inv_order_var(p_k),:);
38         oo.dr.ghxx(oo.dr.inv_order_var(p_a),:)];
39
40 Us2 = [oo.dr.ghuu(oo.dr.inv_order_var(p_k),:);
41         oo.dr.ghuu(oo.dr.inv_order_var(p_a),:)];
42
43 Us3 = [oo.dr.ghxu(oo.dr.inv_order_var(p_k),:);
44         oo.dr.ghxu(oo.dr.inv_order_var(p_a),:)];
45
46 Us0 = [oo.dr.ghs2(oo.dr.inv_order_var(p_k),:);
47         oo.dr.ghs2(oo.dr.inv_order_var(p_a),:)];
48
49 Ux1 = [oo.dr.ghxx(oo.dr.inv_order_var(p_y),:);
50         oo.dr.ghxx(oo.dr.inv_order_var(p_I),:);
51         oo.dr.ghxx(oo.dr.inv_order_var(p_c),:);
52         oo.dr.ghxx(oo.dr.inv_order_var(p_w),:);
53         oo.dr.ghxx(oo.dr.inv_order_var(p_R),:);
54         oo.dr.ghxx(oo.dr.inv_order_var(p_r),:)];
55
56 Ux2 = [oo.dr.ghuu(oo.dr.inv_order_var(p_y),:);
57         oo.dr.ghuu(oo.dr.inv_order_var(p_I),:);
58         oo.dr.ghuu(oo.dr.inv_order_var(p_c),:);
59         oo.dr.ghuu(oo.dr.inv_order_var(p_w),:);
60         oo.dr.ghuu(oo.dr.inv_order_var(p_R),:);
61         oo.dr.ghuu(oo.dr.inv_order_var(p_r),:)];
62
63 Ux3 = [oo.dr.ghxu(oo.dr.inv_order_var(p_y),:);
64         oo.dr.ghxu(oo.dr.inv_order_var(p_I),:);
65         oo.dr.ghxu(oo.dr.inv_order_var(p_c),:);
66         oo.dr.ghxu(oo.dr.inv_order_var(p_w),:);
67         oo.dr.ghxu(oo.dr.inv_order_var(p_R),:);
68         oo.dr.ghxu(oo.dr.inv_order_var(p_r),:)];
69
70 Ux0 = [oo.dr.ghs2(oo.dr.inv_order_var(p_y),:);
71         oo.dr.ghs2(oo.dr.inv_order_var(p_I),:);
72         oo.dr.ghs2(oo.dr.inv_order_var(p_c),:);
73         oo.dr.ghs2(oo.dr.inv_order_var(p_w),:);
74         oo.dr.ghs2(oo.dr.inv_order_var(p_R),:);
75         oo.dr.ghs2(oo.dr.inv_order_var(p_r),:)];

```

To produce impulse responses, here is what I do.

1. I first draw shocks of dimension  $T \times (H1 + H2)$ . I specify  $T = 100$  (I'm average over 100 periods),  $H1 = 50$  (I have a 50 period “burn in”), and  $H2 = 20$  (the dimension of the impulse responses).
2. I simulate out values of the variables  $T$  times using these shocks. So I get  $T$  different simulations with  $H1 + H2$  periods each.

3. I average the simulated series over  $T$  at each horizon.
4. I repeat this process, but add one standard deviation to the shock in period  $H1 + 1$  in generating the series.
5. I take the difference between the second simulation and the first for periods  $H1 + 1$  through  $H2$ . These are the impulse responses.

This will produce impulse responses; these will be close (but not identical to all decimal places) to what Dynare produces. By default, Dynare use  $H1 = 100$  (a 100 period burn-in) and  $T = 50$  (averaging over 50 periods). I'm using different values (even if they were the same, the IRFs wouldn't be identical because we're not drawing the same shocks). Below is my Matlab code to produce impulse responses.

```

1  H1 = 50;
2  H2 = 20;
3  T = 100;
4  % draw shocks
5  randn('seed',666)
6  e = sigmae*randn(T,H1+H2);
7
8  slag = zeros(sn,1); % starting position
9  Ssim = zeros(sn,T,H1+H2);
10 Xsim = zeros(tn-sn,T,H1+H2);
11
12
13 for j = 1:T
14     Ssim(:,j,1) = (1/2)*Us0 + A*slag + B*e(j,1) + (1/2)*Us1*kron(slag,slag)...
15         + (1/2)*Us2*kron(e(j,1),e(j,1)) + Us3*kron(slag,e(j,1));
16     Xsim(:,j,1) = (1/2)*Ux0 + C*slag + D*e(j,1) + (1/2)*Ux1*kron(slag,slag)...
17         + (1/2)*Ux2*kron(e(j,1),e(j,1)) + Ux3*kron(slag,e(j,1));
18
19     for i = 2:H1+H2
20         Ssim(:,j,i) = (1/2)*Us0 + A*Ssim(:,j,i-1) + B*e(j,i) + ...
21             (1/2)*Us1*kron(Ssim(:,j,i-1),Ssim(:,j,i-1)) + ...
22             (1/2)*Us2*kron(e(j,i),e(j,i)) + Us3*kron(Ssim(:,j,i-1),e(j,i));
23         Xsim(:,j,i) = (1/2)*Ux0 + C*Ssim(:,j,i-1) + D*e(j,i) + ...
24             (1/2)*Ux1*kron(Ssim(:,j,i-1),Ssim(:,j,i-1)) + ...
25             (1/2)*Ux2*kron(e(j,i),e(j,i)) + Ux3*kron(Ssim(:,j,i-1),e(j,i));
26     end
27
28 end
29
30 % now calculate means
31 Sm = zeros(sn,H1+H2);
32 Xm = zeros(tn-sn,H1+H2);
33

```



```

34 for k = 1:sn
35     for i = 1:H1+H2
36         Sm(k,i) = mean(Ssim(k,:,i));
37     end
38 end
39
40 for k = 1:tn-sn
41     for i = 1:H1+H2
42         Xm(k,i) = mean(Xsim(k,:,i));
43     end
44 end
45
46 % Now do a second simulation
47 Ssim2 = zeros(sn,T,H1+H2);
48 Xsim2 = zeros(tn-sn,T,H1+H2);
49
50 % Keep e the same, but add sigmae to it in period H1+1
51 e1 = e;
52 e1(:,H1+1) = e1(:,H1+1) + sigmae;
53
54 % Now resimulate using e1
55 for j = 1:T
56     Ssim2(:,j,1) = (1/2)*Us0 + A*slag + B*e1(j,1) + ...
57         (1/2)*Us1*kron(slag,slag) + (1/2)*Us2*kron(e1(j,1),e1(j,1)) ...
58         + Us3*kron(slag,e1(j,1));
59     Xsim2(:,j,1) = (1/2)*Ux0 + C*slag + D*e1(j,1) + ...
60         (1/2)*Ux1*kron(slag,slag) + (1/2)*Ux2*kron(e1(j,1),e1(j,1)) + ...
61         Ux3*kron(slag,e1(j,1));
62
63     for i = 2:H1+H2
64         Ssim2(:,j,i) = (1/2)*Us0 + A*Ssim2(:,j,i-1) + B*e1(j,i) + ...
65             (1/2)*Us1*kron(Ssim2(:,j,i-1),Ssim2(:,j,i-1)) + ...
66             (1/2)*Us2*kron(e1(j,i),e1(j,i)) + ...
67             Us3*kron(Ssim2(:,j,i-1),e1(j,i));
68         Xsim2(:,j,i) = (1/2)*Ux0 + C*Ssim2(:,j,i-1) + D*e1(j,i) + ...
69             (1/2)*Ux1*kron(Ssim2(:,j,i-1),Ssim2(:,j,i-1)) + ...
70             (1/2)*Ux2*kron(e1(j,i),e1(j,i)) + ...
71             Ux3*kron(Ssim2(:,j,i-1),e1(j,i));
72     end
73
74 end
75
76 % now calculate means
77 Sm2 = zeros(sn,H1+H2);
78 Xm2 = zeros(tn-sn,H1+H2);
79
80 for k = 1:sn
81     for i = 1:H1+H2
82         Sm2(k,i) = mean(Ssim2(k,:,i));

```

```

83     end
84 end
85
86 for k = 1:tn-sn
87     for i = 1:H1+H2
88         Xm2(k,i) = mean(Xsim2(k,:,i));
89     end
90 end
91
92 Sirf = zeros(sn,H2);
93 Xirf = zeros(tn-sn,H2);
94
95 for j = 1:H2
96     Sirf(:,j) = Sm2(:,H1+j) - Sm(:,H1+j);
97     Xirf(:,j) = Xm2(:,H1+j) - Xm(:,H1+j);
98 end

```

An irritation sometimes arises here. Unlike in a first order approximation, higher order approximations can explode. In a simple model like this explosion is unlikely for reasonably sized shocks, but you can get the model to be explosive if you make the shocks big enough. In multi-shock models explosion occurs somewhat more frequently.

How do we compute impulse responses and simulations if the coefficients of the second order state space representation are explosive? We can use what is called “pruning.” To get Dynare to do this you just have to added “pruning” as an option in the “stoch\_simul” command. Doing pruning by yourself is fairly straightforward. There are multiple ways of implementing it. The easiest is as follows. When doing the simulation, do both a linear and a non-linear simulation (the linear simulation just uses  $A$ ,  $B$ ,  $C$ , and  $D$  matrixes, which are the same whether you solve the model with a first or a second order approximation). Then, when doing your simulation for the non-linear model, for the “higher order” terms in the approximation (the things multiplied by  $\Upsilon$  coefficients) used the state vector simulated from the linear simulation (but still use the state vector from the non-linear simulation for the linear term). This will guarantee that the simulation does not explode. My code for the pruned construction of impulse responses is given below:

```

1  %%%% Now do this by pruning
2  slag = zeros(sn,1); % starting position
3  Ssim = zeros(sn,T,H1+H2);
4  Xsim = zeros(tn-sn,T,H1+H2);
5  Ssim_lin = zeros(sn,T,H1+H2); % the linear simulation
6
7
8  for j = 1:T
9      Ssim(:,j,1) = (1/2)*Us0 + A*slag + B*e(j,1) + (1/2)*Us1*kron(slag,slag)...
10         + (1/2)*Us2*kron(e(j,1),e(j,1)) + Us3*kron(slag,e(j,1));
11      Ssim_lin(:,j,1) = A*slag + B*e(j,1);
12      Xsim(:,j,1) = (1/2)*Ux0 + C*slag + D*e(j,1) + (1/2)*Ux1*kron(slag,slag)...

```

```

13         + (1/2)*Ux2*kron(e(j,1),e(j,1)) + Ux3*kron(slag,e(j,1));
14
15     for i = 2:H1+H2
16         Ssim(:,j,i) = (1/2)*Us0 + A*Ssim(:,j,i-1) + B*e(j,i) + ...
17             (1/2)*Us1*kron(Ssim_lin(:,j,i-1),Ssim_lin(:,j,i-1)) + ...
18             (1/2)*Us2*kron(e(j,i),e(j,i)) + Us3*kron(Ssim_lin(:,j,i-1),e(j,i));
19         Ssim_lin(:,j,i) = A*Ssim_lin(:,j,i-1) + B*e(j,i);
20         Xsim(:,j,i) = (1/2)*Ux0 + C*Ssim(:,j,i-1) + D*e(j,i) + ...
21             (1/2)*Ux1*kron(Ssim_lin(:,j,i-1),Ssim_lin(:,j,i-1)) + ...
22             (1/2)*Ux2*kron(e(j,i),e(j,i)) + Ux3*kron(Ssim_lin(:,j,i-1),e(j,i));
23     end
24
25 end
26
27 % now calculate means
28 Sm = zeros(sn,H1+H2);
29 Xm = zeros(tn-sn,H1+H2);
30
31 for k = 1:sn
32     for i = 1:H1+H2
33         Sm(k,i) = mean(Ssim(k,:,i));
34     end
35 end
36
37 for k = 1:tn-sn
38     for i = 1:H1+H2
39         Xm(k,i) = mean(Xsim(k,:,i));
40     end
41 end
42
43 % Now do a second simulation
44 Ssim2 = zeros(sn,T,H1+H2);
45 Xsim2 = zeros(tn-sn,T,H1+H2);
46 Ssim2_lin = zeros(sn,T,H1+H2);
47
48 % Keep e the same, but add sigmae to it in period H1+1
49 e1 = e;
50 e1(:,H1+1) = e1(:,H1+1) + sigmae;
51
52 % Now resimulate using e1
53 for j = 1:T
54     Ssim2(:,j,1) = (1/2)*Us0 + A*slag + B*e1(j,1) + ...
55         (1/2)*Us1*kron(slag,slag) + (1/2)*Us2*kron(e1(j,1),e1(j,1)) ...
56         + Us3*kron(slag,e1(j,1));
57     Ssim2_lin(:,j,1) = A*slag + B*e1(j,1);
58     Xsim2(:,j,1) = (1/2)*Ux0 + C*slag + D*e1(j,1) + ...
59         (1/2)*Ux1*kron(slag,slag) + (1/2)*Ux2*kron(e1(j,1),e1(j,1)) + ...
60         Ux3*kron(slag,e1(j,1));
61

```

```

62     for i = 2:H1+H2
63         Ssim2(:,j,i) = (1/2)*Us0 + A*Ssim2(:,j,i-1) + B*e1(j,i) + ...
64             (1/2)*Us1*kron(Ssim2_lin(:,j,i-1),Ssim2_lin(:,j,i-1)) + ...
65             (1/2)*Us2*kron(e1(j,i),e1(j,i)) + ...
66             Us3*kron(Ssim2_lin(:,j,i-1),e1(j,i));
67         Ssim2_lin(:,j,i) = A*Ssim2_lin(:,j,i-1) + B*e1(j,i);
68         Xsim2(:,j,i) = (1/2)*Ux0 + C*Ssim2(:,j,i-1) + D*e1(j,i) + ...
69             (1/2)*Ux1*kron(Ssim2_lin(:,j,i-1),Ssim2_lin(:,j,i-1)) + ...
70             (1/2)*Ux2*kron(e1(j,i),e1(j,i)) + ...
71             Ux3*kron(Ssim2_lin(:,j,i-1),e1(j,i));
72     end
73
74 end
75
76 % now calculate means
77 Sm2 = zeros(sn,H1+H2);
78 Xm2 = zeros(tn-sn,H1+H2);
79
80 for k = 1:sn
81     for i = 1:H1+H2
82         Sm2(k,i) = mean(Ssim2(k,:,i));
83     end
84 end
85
86 for k = 1:tn-sn
87     for i = 1:H1+H2
88         Xm2(k,i) = mean(Xsim2(k,:,i));
89     end
90 end
91
92 Sirf_prune = zeros(sn,H2);
93 Xirf_prune = zeros(tn-sn,H2);
94
95 for j = 1:H2
96     Sirf_prune(:,j) = Sm2(:,H1+j) - Sm(:,H1+j);
97     Xirf_prune(:,j) = Xm2(:,H1+j) - Xm(:,H1+j);
98 end

```

This code is virtually the same as the one I used above. The only differences are that (1) I compute a linear simulation of the state along with the non-linear simulation, and (2) I use the value of the state vector from the linear simulation only in the higher order terms of the non-linear simulation. You can verify that there are only tiny differences between the pruned and standard impulse responses. To see why and where pruning can make a difference, increase the shock standard deviation by 100 – you’ll see that the standard approach explodes and therefore can’t be used to construct impulse responses, whereas the pruned simulation works.

Another area in which a higher order approximation is useful is in doing welfare analysis. Welfare is the present discounted value of household utility. You can include a recursive representation of

welfare in your Dynare code as follows:

$$V_t = \frac{c_t^{1-\sigma} - 1}{1-\sigma} + \beta E_t V_{t+1} \quad (3)$$

This looks like a Bellman equation, but there is no max operator (this is because implicitly in the code  $c_t$  is chosen optimally). Since I'm using  $\sigma = 1$  and Matlab/Dynare can't understand limits, when writing this in my code I need to use  $\ln C_t$  in flow utility. To include this in Dynare, declare  $V$  as a variable and add the following equilibrium condition:

```
1 V = c + beta*V(+1);
```

Note that because my Dynare code interprets  $c$  as  $\ln c_t$ , this is why I just have this showing up as  $c$  (e.g. utility is not linear). This is all I have to do. The inclusion of this variable does not affect the Dynamics of any other variable in the model, it just allows me to look at how  $V$  (e.g. welfare) is affected.

The steady state value of  $V$  is easily seen to be  $\frac{1}{1-\beta} \ln c^*$  – e.g. this is just proportion to the log of steady state consumption. In a first order approximation, the expected values of variables correspond to their steady state values. This would tell you that average welfare is the same in an economy with a standard deviation of productivity shocks of 0.01 versus an economy with a standard deviation of productivity shocks of 0.02. But we know this can't necessarily be right – households have a preference to smooth consumption, so one would think that bigger shocks would make them have lower expected welfare.

We can see this if we use a higher order approximation. First, I solve the model using a second order approximation where the standard deviation of productivity shocks is 0.01. The output from Dynare is found below:

```
Configuring Dynare ...
[mex] Generalized QZ.
[mex] Sylvester equation solution.
[mex] Kronecker products.
[mex] Sparse kronecker products.
[mex] Local state space iteration (second order).
[mex] Bytecode evaluation.
[mex] k-order perturbation solver.
[mex] k-order solution simulation.
[mex] Quasi Monte-Carlo sequence (Sobol).
[mex] Markov Switching SBVAR.
```

```
Starting Dynare (version 4.4.2).
Starting preprocessing of the model file ...
Found 9 equation(s).
```

```

Evaluating expressions...done
Computing static model derivatives:
  - order 1
Computing dynamic model derivatives:
  - order 1
  - order 2
Processing outputs ...done
Preprocessing completed.
Starting MATLAB/Octave computing.

```

#### STEADY-STATE RESULTS:

```

y    1.12546
I    -0.312508
k    3.37637
a    0
c    0.854424
w    0.719993
R    -3.34953
r    0.010101
V    85.4424

```

#### MODEL SUMMARY

```

Number of variables:      9
Number of stochastic shocks: 1
Number of state variables: 2
Number of jumpers:       3
Number of static variables: 5

```

#### MATRIX OF COVARIANCE OF EXOGENOUS SHOCKS

```

Variables      e
e              0.000100

```

#### POLICY AND TRANSITION FUNCTIONS

	y	I	k	a	c
Constant	1.125457	-0.312578	3.376370	0	0.854424

(correction)	0	-0.000070	-0.000002	0	0.0
k(-1)	0.333333	-0.502551	0.962436	0	0.5
a(-1)	0.900000	3.139964	0.078499	0.900000	0.2
e	1.000000	3.488849	0.087221	1.000000	0.2
k(-1),k(-1)	0	-0.536347	0.014107	0	0.0
a(-1),k(-1)	0	2.684454	-0.047889	0	-0.0
a(-1),a(-1)	0	-3.408636	0.034945	0	0.0
e,e	0	-4.208192	0.043142	0	0.0
k(-1),e	0	2.982727	-0.053210	0	-0.0
a(-1),e	0	-7.574746	0.077656	0	0.0

#### APROXIMATED THEORETICAL MOMENTS

VARIABLE	MEAN	STD. DEV.	VARIANCE
y	1.1256	0.0286	0.0008
I	-0.3145	0.0743	0.0055
k	3.3768	0.0275	0.0008
a	-0.0000	0.0229	0.0005
c	0.8547	0.0194	0.0004
w	0.7201	0.0286	0.0008
R	-3.3498	0.0212	0.0005
r	0.0101	0.0007	0.0000
V	85.4704	0.5380	0.2895

#### APPROXIMATED MATRIX OF CORRELATIONS

Variables	y	I	k	a	c	w	R	r	V
y	1.0000	0.9017	0.7567	0.9600	0.8569	1.0000	0.4202	0.3167	0.9517
I	0.9017	1.0000	0.3997	0.9867	0.5499	0.9017	0.7712	0.6957	0.7254
k	0.7567	0.3997	1.0000	0.5433	0.9854	0.7567	-0.2752	-0.3804	0.9209
a	0.9600	0.9867	0.5433	1.0000	0.6783	0.9600	0.6576	0.5697	0.8276
c	0.8569	0.5499	0.9854	0.6783	1.0000	0.8569	-0.1076	-0.2175	0.9738
w	1.0000	0.9017	0.7567	0.9600	0.8569	1.0000	0.4202	0.3167	0.9517
R	0.4202	0.7712	-0.2752	0.6576	-0.1076	0.4202	1.0000	0.9938	0.1213
r	0.3167	0.6957	-0.3804	0.5697	-0.2175	0.3167	0.9938	1.0000	0.0102
V	0.9517	0.7254	0.9209	0.8276	0.9738	0.9517	0.1213	0.0102	1.0000

# APPROXIMATED COEFFICIENTS OF AUTOCORRELATION

Order	1	2	3	4	5
y	0.9367	0.8784	0.8246	0.7748	0.7289
I	0.8827	0.7777	0.6839	0.6001	0.5252
k	0.9980	0.9925	0.9840	0.9730	0.9597
a	0.9000	0.8100	0.7290	0.6561	0.5905
c	0.9927	0.9826	0.9702	0.9559	0.9398
w	0.9367	0.8784	0.8246	0.7748	0.7289
R	0.8768	0.7668	0.6686	0.5811	0.5031
r	0.8868	0.7853	0.6945	0.6133	0.5406
V	0.9746	0.9490	0.9233	0.8975	0.8718
Total computing time : 0h00m02s					

Note that the average value of  $V$  is 85.4704, which is actually higher than the steady state value (which evidently means that you prefer, in an average sense, stochastic productivity shocks to no productivity shocks). This might seem somewhat odd since you have a preference to smooth consumption. I actually have a paper on this with Mike Pries and Rob Lester in the *Journal of Economic Dynamics and Control*. It turns out that it is conceivable that you prefer more shock variance to less depending on parameters, but there is something about which you want to be careful here. What matters for agents in the economy is the level of  $a_t$ . I wrote the process for  $a_t$  in the log. It turns out that the mean of  $a_t$  is increasing in the variance of  $\ln a_t$  via a Jensen's inequality type argument. An alternative way to write the process for  $a_t$  is as an AR(1) in the level with a mean of zero:

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

It turns out that if you do this, a higher variance of  $e_t$  reduces expected welfare. This is because writing the process this way the stochastic mean of  $a_t$  is unaffected by the variance of the shock. When writing the process for  $a_t$  in the log you are confounding two effects – more variance with a higher mean of  $a_t$ . When writing the process this way, you are isolating only the effect of the variance. For more variance to be desirable for welfare, you need factor supply to be very elastic, which it is not here. I bring this subtle point up because we usually don't worry about whether we write the exogenous processes in levels of log-levels. To a first order approximation, they are the same. But to higher order this stuff can matter.

Another subtle point here that emerges is whether one should evaluate welfare conditional on a particular realization of a state or unconditionally. By unconditional welfare I mean the exercise I undertook above – look at the stochastic mean of  $V_t$ . You can compare two different regimes



by doing this (i.e. a high and low variance regime, or in a more involved model different policy regimes). But the comparisons are in a sense not fair because the different regimes fluctuate around different stochastic means. The example I gave above is an extreme example of this. An alternative is to look at conditional welfare. For example, you can condition on looking at the non-stochastic steady state, and evaluate welfare in two different regimes at that point (rather than looking at unconditional welfare).

There is increasing interest in the effects of risk or uncertainty shocks – in essence, shocks to the variance of shocks. These shocks obviously have no effect in a first order approximation. Interesting, they also have no (important) effect in a second order approximation (I say important because they can affect means but do not affect dynamics). The basic intuition for this is based on studying risk aversion and precautionary saving – the second derivative of utility being negative says that people don't like risk, but whether or not they act on it depends on the third derivative of utility being non-zero (if it is positive, they respond to more uncertainty by trying to save). Hence, to get large effects of uncertainty shocks, you need at least a third order approximation. I will not discuss how to do that here, but Dynare is capable of doing it.