# Solving DSGE models Macro II - Fluctuations - ENSAE, 2024-2025

Pablo Winant

2025-03-12

Introduction

What is the main specificity of economic modeling? In (macro)economics, we *model* the behaviour of economic agents by specifying:

their objective

$$\max_{c_t} E_t \sum_{s \geq t} \beta^s U(c_s)$$



 $\max \pi_t$ ...

their constraints (budget constraint, econ. environment...) What is the main specificity of economic modeling? In (macro)economics, we *model* the behaviour of economic agents by specifying:

their objective

$$\max_{c_t} E_t \sum_{s \geq t} \beta^s U(c_s)$$

 $\max \pi_{t}$ 



their constraints (budget constraint, econ. environment...)

This has important implications:

macro models are *forward looking* rely on expectations
 macro models need to be **solved**

In many cases, there is no closed form for the solution -> we need numerical techniques

 1996: Michel Juillard created an opensource software to solve DSGE models



Figure 1: Michel Juillard

It has been widely adopted:
 early version in Gauss
 then Matlab/Octave/Scilab
 latest version in Julia
 ... and Python (checkout dyno )

- 1996: Michel Juillard created an opensource software to solve DSGE models
  - DSGE: Dynamic Stochastic General Equilibrium



Figure 1: Michel Juillard

It has been widely adopted:
 early version in Gauss
 then Matlab/Octave/Scilab
 latest version in Julia
 ... and Python (checkout dyno )

- 1996: Michel Juillard created an opensource software to solve DSGE models
  - DSGE: Dynamic Stochastic General Equilibrium
  - usually solved around a steady-state



Figure 1: Michel Juillard

It has been widely adopted:
 early version in Gauss
 then Matlab/Octave/Scilab
 latest version in Julia
 ... and Python (checkout dyno )

- 1996: Michel Juillard created an opensource software to solve DSGE models
  - DSGE: Dynamic Stochastic General Equilibrium
  - usually solved around a steady-state
  - Now about 10 contributors.
- It has been widely adopted:
   early version in Gauss
   then Matlab/Octave/Scilab
   latest version in Julia
   ... and Python (checkout dyno )



Figure 1: Michel Juillard

- 1996: Michel Juillard created an opensource software to solve DSGE models
  - DSGE: Dynamic Stochastic General Equilibrium
  - usually solved around a steady-state
- Now about 10 contributors.
  - + power users who have contributed to the code
- It has been widely adopted:
  - early version in Gauss
  - then Matlab/Octave/Scilab
  - latest version in Julia
  - ... and Python (checkout dyno )



Figure 1: Michel Juillard

# DSGE Models in institutions

Nowadays most DSGE models built in institutions have a Dynare version (IMF/GIMF, EC/Quest, ECB/, NYFed/FRBNY)

- they are usually based on the *midsize model* from Smets & Wouters (10 equations)
- but have grown up a lot (»100 equations)

# DSGE Models in institutions

Nowadays most DSGE models built in institutions have a Dynare version (IMF/GIMF, EC/Quest, ECB/, NYFed/FRBNY)

- they are usually based on the *midsize model* from Smets & Wouters (10 equations)
- but have grown up a lot (»100 equations)

Institutions, led by researchers are diversifying their model

- Semi-Structural Models
- Computational General Equilibrium Models
- Network Models
- Agent-based Models
- Heterogenous Agents Models

# The Plan

Provide a short introduction to DSGE modeling:

- How models are solved (today)
- Small Open Economy (aka IRBC model)
- Heterogeneity
- Financial Intermediation

In passing, we'll discuss some of the trends

# Solving a model

# Model

A very concise representation of a model

$$\mathbb{E}_t\left[f(y_{t+1},y_t,y_{t-1},\epsilon_t)\right]=0$$

The problem:

y<sub>t</sub> ∈ ℝ<sup>n</sup>: the vector of endogenous variables
 ϵ<sub>t</sub> ∈ ℝ<sup>n<sub>e</sub></sup>: the vector of exogenous variables
 we assume that ϵ<sub>t</sub> is a zero-mean gaussian process
 f : ℝ<sup>n</sup> → ℝ<sup>n</sup>: the model equations

The **solution**:

g such that

$$\forall t, y_t = g(y_{t-1}, \epsilon_t)$$

# The timing of the equations

🖣 Tip

In a dynare modefile the model equations are coded in the model; ...; end; block. Variable  $v_t$  (resp  $v_{t-1}$ ,  $v_{t+1}$ ) is denoted by v or v(0) (resp v(-1), v(+1)).

#### **General Timing Convention**

New information arrives with the innovations  $\epsilon_t$ .

At date t, the information set is spanned by  $\mathcal{F}_t=\mathcal{F}(\cdots,\epsilon_{t-3},\epsilon_{t-2},\epsilon_{t-1},\epsilon_t)$ 

By convention an endogenous variable has a subscript t if it is known first at date t.

# The timing of the equations

🖣 Tip

In a dynare modefile the model equations are coded in the model; ...; end; block. Variable  $v_t$  (resp  $v_{t-1}$ ,  $v_{t+1}$ ) is denoted by v or v(0) (resp v(-1), v(+1)).

#### **General Timing Convention**

New information arrives with the innovations  $\epsilon_t$ .

At date t, the information set is spanned by  $\mathcal{F}_t=\mathcal{F}(\cdots,\epsilon_{t-3},\epsilon_{t-2},\epsilon_{t-1},\epsilon_t)$ 

By convention an endogenous variable has a subscript t if it is known first at date t.

Several **variable types** depending on how they appear in the model:

**N 1 1 1 1 1 1 1** 

# The timing of equations

#### Example

Using Dynare's timing conventions:

- Write the production function in the RBC
- Write the law of motion for capital k, with a depreciation rate
  - $\delta$  and investment i
    - when is capital known?
    - when is investment known?
- Add a multiplicative investment efficiency shock  $\chi_t$ . Assume it is an AR1 driven by innovation  $\eta_t$  and autocorrelation  $\rho_{\gamma}$ 
  - how do you write the law of motion for capital?

### Steady-state

The deterministic steady-state satisfies:

$$f(\overline{y},\overline{y},\overline{y},\overline{y},0)=0$$

Often, there is a closed-form solution.

Otherwise, one must resort to a numerical solver to solve

$$\overline{y} \to f(\overline{y},\overline{y},\overline{y},0)$$

#### 🖣 Tip

In dynare the steady-state values are provided in the steadystate\_model; ...; end; block. One can check they are correct using the check; statement. To find numerically the steady-state: steady;.

## The implicit system

Replacing the solution

$$y_t = g(y_{t-1}, \epsilon_t)$$

in the system

$$\mathbb{E}_t\left[f(y_{t+1},y_t,y_{t-1},\epsilon_t)\right]=0$$

we obtain:

 $\mathbb{E}_t\left[f(g(g(y_{t-1},\epsilon_t),\epsilon_{t+1}),g(y_{t-1},\epsilon_t),y_{t-1},\epsilon_t)\right]=0$ 

It is an equation defining implicitly the function g()

#### The state-space

 $\mathbb{E}_t\left[f(g(g(y_{t-1},\epsilon_t),\epsilon_{t+1}),g(y_{t-1},\epsilon_t),y_{t-1},\epsilon_t)\right]=0$ 

In this expression,  $y_{t-1}, \epsilon_t$  is the state-space:

 $\blacktriangleright$  it contains all information available at t to predict the future evolution of  $(y_s)_{s\geq t}$ 

#### The state-space

 $\mathbb{E}_t\left[f(g(g(y_{t-1},\epsilon_t),\epsilon_{t+1}),g(y_{t-1},\epsilon_t),y_{t-1},\epsilon_t)\right]=0$ 

In this expression,  $y_{t-1}, \epsilon_t$  is the state-space:

 $\blacktriangleright$  it contains all information available at t to predict the future evolution of  $(y_s)_{s\geq t}$ 

Dropping the time subscripts, the equation must be satisfied for any realization of  $(\boldsymbol{y}, \boldsymbol{\epsilon})$ 

 $\forall (y,\epsilon) \ \Phi(g)(y,\epsilon) = \mathbb{E}_{\epsilon'} \left[ f(g(g(y,\epsilon),\epsilon'),g(y,\epsilon),y,\epsilon) \right] = 0$ 

It is a functional equation  $\Phi(g) = 0$ 

## Expected shocks

=

 $\approx$ 

First order approximation:

Assume 
$$|\epsilon| << 1, |\epsilon'| << 1$$

Perform a Taylor expansion with respect to future shock:

$$\mathbb{E}_{\epsilon'}\left[f(g(g(y,\epsilon),\epsilon'),g(y,\epsilon),y,\epsilon)\right] \quad (1)$$

 $\mathbb{E}_{\epsilon'}\left[f(g(g(y,\epsilon),0),g(y,\epsilon),y,\epsilon)\right]$  (2)

$$+\mathbb{E}_{\epsilon'}\left[f'_{y_{t+1}}(g(g(y,\epsilon),0),g(y,\epsilon),y,\epsilon)g'_{\epsilon}\epsilon'\right]+o(\epsilon')$$
(3)

$$f(g(g(y,\epsilon),0),g(y,\epsilon),y,\epsilon)$$
 (4)

## Expected shocks

First order approximation:

Assume 
$$|\epsilon| << 1, |\epsilon'| << 1$$

Perform a Taylor expansion with respect to future shock:

$$\begin{split} & \mathbb{E}_{\epsilon'}\left[f(g(g(y,\epsilon),\epsilon'),g(y,\epsilon),y,\epsilon)\right] & (1) \\ = & \mathbb{E}_{\epsilon'}\left[f(g(g(y,\epsilon),0),g(y,\epsilon),y,\epsilon)\right] & (2) \\ & +\mathbb{E}_{\epsilon'}\left[f'_{y_{t+1}}(g(g(y,\epsilon),0),g(y,\epsilon),y,\epsilon)g'_{\epsilon}\epsilon'\right] + o(\epsilon') & (3) \\ \approx & f(g(g(y,\epsilon),0),g(y,\epsilon),y,\epsilon) & (4) \end{split}$$

This uses the fact that  $\mathbb{E}[\epsilon'] = 0$ .

At first order, expected shocks play no role.

To capture precautionary behaviour (like risk premia), we would need to increase the approximation order.

#### First order perturbation

We are left with the system:

$$F(y,\epsilon) = f(g(g(y,\epsilon),0),g(y,\epsilon),y,\epsilon) = 0$$

A variant of the *implicit function theorem* then yields the existence of a first approximation of g:

$$g(y,\epsilon)=\overline{y}+g_y'(y-\overline{y})+g_e'\epsilon_t$$

#### First order perturbation

We are left with the system:

$$F(y,\epsilon) = f(g(g(y,\epsilon),0),g(y,\epsilon),y,\epsilon) = 0$$

A variant of the *implicit function theorem* then yields the existence of a first approximation of g:

$$g(y,\epsilon)=\overline{y}+g_y'(y-\overline{y})+g_e'\epsilon_t$$

Unknown quantities  $g'_y$ , and  $g'_e$  are obtained using the *method of undeterminate coefficients*. Plug the first approximation into the system and write the conditions

$$F'_{y}(\overline{y}, 0) = 0$$
$$F'_{\epsilon}(\overline{y}, 0) = 0$$

# Computing $g_y^{'}$

Recall the system:

$$F(y,\epsilon)=f(g(g(y,\epsilon),0),g(y,\epsilon),y,\epsilon)=0$$

We have

$$F'_y(\overline{y},0) = f'_{y_{t+1}}g'_yg'_y + f'_{y_t}g'_y + f'_{y_{t-1}} = 0$$

# Computing $g_{y}^{'}$

Recall the system:

$$F(y,\epsilon)=f(g(g(y,\epsilon),0),g(y,\epsilon),y,\epsilon)=0$$

We have

$$F'_y(\overline{y},0) = f'_{y_{t+1}}g'_yg'_y + f'_{y_t}g'_y + f'_{y_{t-1}} = 0$$

 $g'_{y}$  is the solution of a specific Riccatti equation

$$AX^2 + BX + C$$

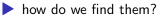
where A, B, C and  $X = g'_y$  are square matrices  $\in \mathbb{R}^n \times \mathbb{R}^n$ 

## First Order Deterministic Model

Let's pause a minute to observe the first order deterministic model:

 $AX^2 + BX + C$ 

From our intuition in dimension 1, we know there must be multiple solutions



how do we select the right ones?

I the absence of shocks the dynamics of the model are given by

$$y_t = Xy_{t-1}$$

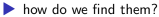
What is the condition for the model to be stationary?

## First Order Deterministic Model

Let's pause a minute to observe the first order deterministic model:

 $AX^2 + BX + C$ 

From our intuition in dimension 1, we know there must be multiple solutions



how do we select the right ones?

I the absence of shocks the dynamics of the model are given by

$$y_t = Xy_{t-1}$$

What is the condition for the model to be stationary?

-> the biggest eigenvalue of X should be smaller than 1

# Multiplicity of solution

It is possible to show that the system is associated with 2n generalized eigenvalues:

$$|\lambda_1| \leq \cdots \leq |\lambda_{2n}|$$

For each choice C of n eigenvalues (|C| = n), a specific recursive solution  $X_C$  can be *constructed*. It has eigenvalues C.

# Multiplicity of solution

It is possible to show that the system is associated with 2n generalized eigenvalues:

$$|\lambda_1| \leq \cdots \leq |\lambda_{2n}|$$

For each choice C of n eigenvalues (|C| = n), a specific recursive solution  $X_C$  can be *constructed*. It has eigenvalues C.

This yields at least  $\binom{2n}{n}$  different combinations.

# Multiplicity of solution

It is possible to show that the system is associated with 2n generalized eigenvalues:

$$|\lambda_1| \leq \cdots \leq |\lambda_{2n}|$$

For each choice C of n eigenvalues (|C| = n), a specific recursive solution  $X_C$  can be *constructed*. It has eigenvalues C.

This yields at least  $\binom{2n}{n}$  different combinations.

A model is well defined when there is **exactly one solution that is non divergent**.

This is equivalent to:

$$|\lambda_1| \leq \cdots \leq |\lambda_n| \leq 1 < |\lambda_{n+1}| \leq \cdots \leq |\lambda_{2n}|$$

Forward looking inflation:

$$\pi_t = \alpha \pi_{t+1}$$

with  $\alpha < 1$ .

Is it well defined?

Forward looking inflation:

$$\pi_t = \alpha \pi_{t+1}$$

with  $\alpha < 1$ .

Is it well defined?

We can rewrite the system as:

$$\alpha \pi_{t+1} - \pi_t + 0\pi_{t-1} = 0$$

or

$$\pi_{t+1} - \left(\frac{1}{\alpha} + 0\right)\pi_t + \left(\frac{1}{\alpha}0\right)\pi_{t-1} = 0$$

Forward looking inflation:

$$\pi_t = \alpha \pi_{t+1}$$

with  $\alpha < 1$ .

Is it well defined?

We can rewrite the system as:

$$\alpha \pi_{t+1} - \pi_t + 0\pi_{t-1} = 0$$

or

$$\pi_{t+1} - \left(\frac{1}{\alpha} + 0\right)\pi_t + \left(\frac{1}{\alpha}0\right)\pi_{t-1} = 0$$

The generalized eigenvalues are  $0 \le 1 < \frac{1}{\alpha}$ .

Forward looking inflation:

$$\pi_t = \alpha \pi_{t+1}$$

with  $\alpha < 1$ .

Is it well defined?

We can rewrite the system as:

$$\alpha \pi_{t+1} - \pi_t + 0\pi_{t-1} = 0$$

or

$$\pi_{t+1} - \left(\frac{1}{\alpha} + 0\right)\pi_t + \left(\frac{1}{\alpha}0\right)\pi_{t-1} = 0$$

The generalized eigenvalues are  $0 \leq 1 < \frac{1}{\alpha}.$ 

The unique stable solution is  $\pi_{4} = 0\pi_{4-1}$ 

Debt accumulation equation by a rational agent:

$$b_{t+1} - (1+\frac{1}{\beta})b_t + \frac{1}{\beta}b_{t-1} = 0$$

Is it well-defined?

Debt accumulation equation by a rational agent:

$$b_{t+1} - (1+\frac{1}{\beta})b_t + \frac{1}{\beta}b_{t-1} = 0$$

Is it well-defined?

Two generalized eigenvalues  $\lambda_1=1<\lambda_2=\frac{1}{\beta}$ 

Debt accumulation equation by a rational agent:

$$b_{t+1} - (1+\frac{1}{\beta})b_t + \frac{1}{\beta}b_{t-1} = 0$$

Is it well-defined?

Two generalized eigenvalues  $\lambda_1 = 1 < \lambda_2 = \frac{1}{\beta}$ 

The unique non-diverging solution is  $b_t = b_{t-1}$ .

it is a unit-root: any initial deviation in b<sub>t-1</sub> has persistent effects

Productivity process:

$$z_t = \rho z_{t-1}$$

with  $\rho < 1:$  well defined

Productivity process:

$$z_t = \rho z_{t-1}$$

with  $\rho < 1$ : well defined

In that case there is a hidden infinite eigenvalue  $\infty$  associated to  $z_{t+1}.$ 

Productivity process:

$$z_t = \rho z_{t-1}$$

with  $\rho < 1:$  well defined

In that case there is a hidden infinite eigenvalue  $\infty$  associated to  $z_{t+1}.$ 

To see why consider the system associated with eigenvalues m and  $\rho {:}$ 

$$z_{t+1}-(m+\rho)z_t+m\rho z_{t-1}=0$$

$$\frac{1}{m}z_{t+1} - (1+\frac{\rho}{m})z_t + \rho z_{t-1} = 0$$

Which corresponds to the initial model when  $m=\infty$ 

Productivity process:

$$z_t = \rho z_{t-1}$$

with  $\rho < 1:$  well defined

In that case there is a hidden infinite eigenvalue  $\infty$  associated to  $z_{t+1}.$ 

To see why consider the system associated with eigenvalues m and  $\rho {\rm :}$ 

$$z_{t+1}-(m+\rho)z_t+m\rho z_{t-1}=0$$

$$\frac{1}{m}z_{t+1} - (1+\frac{\rho}{m})z_t + \rho z_{t-1} = 0$$

Which corresponds to the initial model when  $m=\infty$ 

The generalized eigenvalues are  $\lambda_1=\rho\leq 1<\lambda_2=\infty$ 

More generally, any variable that does not appear in t + 1 creates one infinite generalized eigenvalue.

#### A criterium for well-definedness

Looking again at the list of eigenvalues we set aside the infinite ones.

The model is well specified iff we can sort the eigenvalues as:

$$|\lambda_1| \leq \cdots \leq |\lambda_n| \leq 1 < |\lambda_{n+1}| \leq \cdots |\lambda_{n+k}| \leq \underbrace{|\lambda_{n+k+1}| \cdots \leq |\lambda_{2n}|}_{\text{infinite eigenvalues}}$$

#### i Blanchard-Kahn criterium

The model satisfies the Blanchard-Kahn criterium if the number of eigenvalues greater than one, is exactly equal to the number of variables *appearing* in t + 1. In that case the model is well-defined.

## Computing the solution

There are several classical methods to compute the solution to the algebraic Riccatti equation:

$$AX^2 + BX + C = 0$$

qz decomposition
 traditionnally used in the DSGE literature since Chris Sims
 a little bit unintuitive
 cyclic reduction

 new default in dynare, more adequate for big models
 linear time iteration cf @sec:linear\_time\_iteration
 conceptually very simple

# Computing $g_{e}^{'}$

Now we have  $g_y^\prime,$  how do we get  $g_e^\prime?$ 

Recall:

$$F(y,\epsilon)=f(g(g(y,\epsilon),0),g(y,\epsilon),y,\epsilon)=0$$

We have

$$F'_e(\overline{y},0) = f'_{y_{t+1}}g'_yg'_e + f'_{y_t}g'_e + f'_{\epsilon_t} = 0$$

Now this is easy:

$$g'_e = -(f'_{y_{t+1}}g'_y + f'_{y_t})^{-1}f'_{\epsilon_t} = 0$$

#### The model solution

The result of the model solution:

$$y_t = g_y y_{t-1} + g_e \epsilon_t$$

It is an AR1, driven by exogenous shock  $\epsilon_t$ .

#### The model solution

The result of the model solution:

$$y_t = g_y y_{t-1} + g_e \epsilon_t$$

It is an AR1, driven by exogenous shock  $\epsilon_t$ .

Because it is a well known structure, one can investigate the model with

- impulse response functions
- stochastic simulations

#### The model solution

The result of the model solution:

$$y_t = g_y y_{t-1} + g_e \epsilon_t$$

It is an AR1, driven by exogenous shock  $\epsilon_t$ .

Because it is a well known structure, one can investigate the model with

- impulse response functions
- stochastic simulations

Then to compare the model to the data we compute

implied moments:
 covariances, autocorrelation
 likelihood

Optimizing the fit to the data is called *model* estimation

## Conclusion

#### What can you do with the solution

The solution of a model found by Dynare has an especially simple form: an AR1

$$\blacktriangleright y_t = Xy_{t-1} + Y\epsilon_t$$

 $\blacktriangleright$  where the covariances  $\Sigma$  of  $\epsilon_t$  can be chosen by the modeler

#### What can you do with the solution

The solution of a model found by Dynare has an especially simple form: an AR1

 $\downarrow y_t = Xy_{t-1} + Y\epsilon_t$ 

 $\blacktriangleright$  where the covariances  $\Sigma$  of  $\epsilon_t$  can be chosen by the modeler

With this solution we can (cf next TD)

compute (conditional and unconditional) moments perform stochastic simulations, impulse response function

#### What can you do with the solution

The solution of a model found by Dynare has an especially simple form: an AR1

 $\downarrow y_t = Xy_{t-1} + Y\epsilon_t$ 

 $\blacktriangleright$  where the covariances  $\Sigma$  of  $\epsilon_t$  can be chosen by the modeler

With this solution we can (cf next TD)

compute (conditional and unconditional) moments perform stochastic simulations, impulse response function

# Going Further

Taking the model to the data with Dynare



"estimate" the model: compute the likelihood of a solution and maximize it by choosing the right parameters

"identify" shocks in the data

#### Other functions

- higher order approximation
- (noninear) perfect foresight simulations
- ramsey plan
- discretionary policy
- **•** ...

#### Coming Next



Many models

#### Appendix: Linear Time Iteration

#### Linear Time Iteration

Recall the system to solve:

$$F(y,\epsilon)=f(g(g(y,\epsilon),0),g(y,\epsilon),y,\epsilon)=0$$

but now assume the decision rules today and tomorrow are different:

$$\begin{array}{l} \blacktriangleright \text{ today: } y_t = g(y_{t-1},\epsilon_t) = \overline{y} + Xy_{t-1} + g_y \epsilon_t \\ \blacktriangleright \text{ tomorrow: } y_{t+1} = \widetilde{g}(y_t,\epsilon_{t+1}) = \overline{y} + \widetilde{X}y_{t-1} + \widetilde{g}_y \epsilon_t \end{array}$$

Then the Ricatti equation is written:

$$A\tilde{X}X + BX + C = 0$$

# Linear Time Iteration (2)

The linear time iteration algorithm consists in solving the decision rule X today as a function of decision rule tomorrow  $\tilde{X}$ .

This corresponds to the simple formula:

$$X=-(A\tilde{X}+B)^{-1}C$$

And the full algorithm can be described as:

## Linear Time Iteration (3)

It can be shown that, starting from a random initial guess, the linear time-iteration algorithm converges to the solution X with the smallest modulus:

$$\underbrace{|\lambda_1| \leq \cdots \leq |\lambda_n|}_{\text{Selected eigenvalues}} \leq |\lambda_{n+1}| \cdots \leq |\lambda_{2n}|$$

In other words, it finds the right solution when the model is well specified.

How do you check it is well specified?

$$\begin{array}{l} \lambda_n \text{ is the biggest eigenvalue of solution } X \\ \hline \text{ what about } \lambda_{n+1}? \\ \hline \frac{1}{\lambda_{n+1}} \text{ is the biggest eigenvalue of } (AX+B)^{-1}A \end{array}$$

## Linear Time Iteration (4)

Define

$$M(\lambda) = A\lambda^2 + B\lambda + C$$

For any solution  $X,\,M(\lambda)$  can be factorized as:  $^1$ 

$$M(\lambda) = (\lambda A + AX + B)(\lambda I - X)$$

and

$$det(M(\lambda)) = \underbrace{det(\lambda A + AX + B)}_{Q(\lambda)} det(\lambda I - X)$$

By construction  $Q(\lambda)$  is a polynomial whose roots are those that are not selected by the solution i.e.  $\Lambda \setminus Sp(X)$ .

<sup>&</sup>lt;sup>1</sup>Special case of Bezout theorem. Easy to check in that case

#### Linear Time Iteration (5)

For  $\lambda \neq 0$  we have:

$$\begin{split} \lambda &\in Sp((AX+B)^{-1}A) \\ \Leftrightarrow \ det((AX+B)^{-1})A - I\lambda) = 0 \\ \Leftrightarrow \ det(\frac{1}{\lambda}A - I(AX+B)) = 0 \\ \Leftrightarrow \ Q(\frac{1}{\lambda}) = 0 \\ \Leftrightarrow \ \frac{1}{\lambda} \in G \smallsetminus Sp(X) \end{split}$$

In words,  $(AX+B)^{-1}$  contains all the eigenvalues that have been rejected by the selection of  $X. \label{eq:alpha}$ 

In particular,  $\rho((AX+B)^{-1})A)=1/\min(G\smallsetminus Sp(X))$