# **Exploiting Symmetry in High-Dimensional Dynamic Programming**

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#### **Motivation**

- Most dynamic models in macro (and other fields) deal with either:
  - Representative agent or few agents.
  - A continuum of agents.
- However, many models of interest in macro (IO and trade) deal with finite (but large) number of agents and idiosyncratic/aggregate uncertainty:
  - Industry dynamics with many firms, agents and industries, even models with networks.
  - Heterogeneous agent labor models (e.g., overlapping generations, different types).
- These models are becoming increasingly popular, but:
  - They pose computational challenges as we add more agents.
  - No (non-heuristic) algorithm exists providing global solutions in the presence of aggregate uncertainty.

## Challenges: the curse of dimensionality in equilibrium models

Three components to the curse of dimensionality with many agents (Bellman, 1958, p. IX)

- 1. The cardinality of the state space is enormous.
  - With 266 state variables, with 2 values per state (zero and one), we have more arrangements (2<sup>266</sup>) than the estimated number of protons in the universe.
- 2. With idiosyncratic and aggregate shocks we need to calculate high-dimensional conditional expectations.
- 3. Finding equilibrium paths to the steady-state (ergodic distributions) are extremely hard in high-dimensions.

#### Contribution I

Inspired by economic theory, providing novel method for **globally** solving high-dimensional heterogeneous agent models with **aggregate shocks** which relies on:

- A symmetry present in many heterogeneous agent models, i.e., exchangeability of agents.
  - Example: In general equilibrium models the Walrasian auctioneer removes indices.
  - The solution (function) must be faithful to this symmetry caused by this exchangeability of agents.

#### **Contribution II**

- Concentration of measures, something that resembles the law of large numbers to deal with conditional expectations (very fast).
  - More agents makes it easier to forecast the evolution of distributions.
  - Conditional on the aggregate shock, we can use something similar law of large numbers to calculate expectations.
- We show how to implement the symmetry when using deep neural networks.

With these we globally solve a model with 10,000 agents which was not possible before.

#### Literature Review

- Deep learning as a functional approximation: Maliar et al. (2019), Fernández-Villaverde et al. (2022), Duarte (2018), Azinovic et al. (2022), Han et al. (2021) (a mean-field approach).
- Symmetry in statistics and machine learning: Bloem-Reddy and Teh (2020), Zaheer et al. (2017), and Yarotsky (2018).
- Symmetry in computer science (MDP/RL): Ravindran and Barto (2001) and Narayanamurthy and Ravindran (2008), van der Pol et al. (2020).
- Symmetry in micro and games: Jovanovic and Rosenthal (1988), Hartford et al. (2016)

# Background: Deep learning for functional equations

## **Equilibrium conditions as functional equations**

Most theoretical models in economics with equilibrium conditions can be written as functional equations:

- Take some function(s)  $\psi \in \Psi$  where  $\psi : \mathcal{X} \to \mathcal{Y}$  (e.g. asset price, investment choice, best-response).
- ullet Domain  ${\mathcal X}$  could be state (e.g. dividends, capital, opponents state) or time if sequential.
- The "model" is  $\ell: \Psi \times \mathcal{X} \to \mathcal{R}$  (e.g., Euler and Bellman residuals, equilibrium FOCs).
- The solution is the root of the model (residuals operator), i.e.,  $\mathbf{0} \in \mathcal{R}$ , at each  $x \in \mathcal{X}$ .

Then a solution is a  $\psi^* \in \Psi$  where  $\ell(\psi^*, x) = 0$  for all  $x \in \mathcal{X}$ . How do we find an approximate solution?

## Classical solution method for functional equations

#### Quick review of collocation-like methods:

- 1. Pick finite set of D points  $\hat{\mathcal{X}} \subset \mathcal{X}$  (e.g., a grid).
- 2. Choose approximation  $\hat{\psi}(\cdot; \theta) \in \mathcal{H}(\Theta)$  with coefficients  $\Theta \subseteq \mathbb{R}^M$  (e.g., Chebyshev polynomials).
- 3. Fit with nonlinear least-squares

$$\min_{\theta \in \Theta} \sum_{\mathbf{x} \in \hat{\mathcal{X}}} \ell(\hat{\psi}(\cdot; \theta), \mathbf{x})^2$$

If  $\theta \in \Theta$  is such that  $\ell(\hat{\psi}(\cdot; \theta), x) = 0$  for all  $x \in \hat{\mathcal{X}}$  we say  $\hat{\psi}(\cdot; \theta)$  interpolates  $\hat{\mathcal{X}}$ .

- 4. The goal is to have good **generalization**:
  - The approximate function is close to the solution outside of  $\hat{\mathcal{X}}$ .

## **Deep Neural Networks**

**Deep learning** is **highly-overparameterized**  $\mathcal{H}(\Theta)$  ( $M \gg D$ ) designed for good generalization.

• Example: one layer neural network,  $\hat{\psi}: \mathbb{R}^Q \to \mathbb{R}$ :

$$\hat{\psi}(x;\theta) = W_2 \cdot \sigma (W_1 \cdot x + b_1) + b_2$$

- $W_1 \in \mathbb{R}^{P \times Q}$ ,  $b_1 \in \mathbb{R}^{P \times 1}$ ,  $W_2 \in \mathbb{R}^{1 \times P}$ , and  $b_2 \in \mathbb{R}$ .
- $\sigma(\cdot)$  is a nonlinear function applied element-wise (e.g.,  $\max\{\cdot,0\}$ ).
- $\Theta \equiv \{b_1, W_1, b_2, W_2\}$  are the coefficients, in this example M = PQ + P + P + 1.
- Making it "deeper" by adding another "layer":  $\hat{\psi}(x;\theta) \equiv W_3 \cdot \sigma(W_2 \cdot \sigma(W_1 \cdot x + b_1) + b_2) + b_3$ .
- Very flexible to design  $\mathcal{H}(\Theta)$  using economic insights (e.g., encode symmetry).
- Composing  $\mathcal{H}(\Theta)$  from multiple function (e.g., deeper) tend to **generalize better** better in practice.

## Over-parameterization and convergence

If the number of coefficients is much larger than the number of grid points  $M \gg D$ , there are many different sets of coefficients that achieve interpolation.

- What is going on?
  - Deep neural networks and their optimizers have an inherent implicit bias toward a unique class of interpolating solutions.
  - Figuring out this property is a very active field in computer science and optimization theory.
  - Converges to "simple" (flat) interpolating functions.
    - They have a built-in Occam's razor.
- Modern ML, uses high-dimensional non-convex optimizations. Does the initialization of the coefficients matter?

## **Application**

## How do we pick our application to show how all this works?

- In terms of application, there are two routes:
  - 1. Introducing a sophisticated application where the method "shines".
  - 2. Or, applying it to a well-known example.
- If I tell you about a sophisticated application, how do we know our "solution" method works?
- So we study a well-known example (with a twist).
- Study the more sophisticated applications in future projects.

## **Our application**

A variation of the Lucas and Prescott (1971) model of investment under uncertainty with N firms.

Why?

- 1. Ljungqvist and Sargent (2018), pp. 226-228, use it to introduce recursive competitive equilibria.
- 2. Simple model that fits in one slide.
- 3. Under one parameterization, the model has a known Linear-Quadratic (LQ) solution, which gives us an exact benchmark.
- 4. By changing one parameter, the model is nonlinear, with no known solution. Our method handles the nonlinear case as easily as the LQ case with high accuracy.

## Investment under uncertainty

- Industry consisting of N > 1 firms, each producing the same good.
- Firm of interest produces output x (x units of capital).
- Thus, the vector  $X \equiv [X_1, \dots X_N]^{\top}$  is the production (or capital) of the whole industry.
- The inverse demand function for the industry is, for some  $\nu \geq 1$  (this is our twist):

$$p(X) = 1 - \frac{1}{N} \sum_{i=1}^{N} X_i^{\nu}$$

- The firm does not consider the impact of its individual decisions on p(X).
- Due to adjustment frictions, investing u has a cost  $\frac{\gamma}{2}u^2$ .
- Law of motion for capital  $x' = (1 \delta)x + u + \sigma w + \eta \omega$  where  $w \sim \mathcal{N}(0, 1)$  an i.i.d. idiosyncratic shock, and  $\omega \sim \mathcal{N}(0, 1)$  an i.i.d. aggregate shock, common to all firms.
- The firm chooses u to maximize  $\mathbb{E}\left[\sum_{t=0}^{\infty} \beta^t \left(p(X)x \frac{\gamma}{2}u^2\right)\right]$ .

### Recursive problem

The recursive problem of the firm taking the exogenous policy  $\hat{u}(\cdot, X)$  for all other firms as given is:

$$v(x,X) = \max_{u} \left\{ p(X)x - \frac{\gamma}{2}u^2 + \beta \mathbb{E}\left[v(x',X')\right] \right\}$$
s.t.  $x' = (1-\delta)x + u + \sigma w + \eta \omega$ 

$$X'_{i} = (1-\delta)X_{i} + \hat{u}(X_{i},X) + \sigma W_{i} + \eta \omega, \quad \text{for } i \in \{1,...,N\}$$

Take FOCs and equation using standard steps to write equilibrium as the LOM and Euler equation

$$\gamma u(x,X) = \beta \mathbb{E} \left[ p(X') + \gamma (1-\delta) u(x',X') \right]$$

## Curse of dimensionality: a closer look

$$\gamma u(x,X) = \beta \mathbb{E} \left[ p(X') + \gamma (1-\delta) u(x',X') \right]$$

Let's say there are 1000 agents:

- 1. The domain of the unknown functions of interest  $u(\cdot, \cdot)$  is 1000-dimensional.
- 2. The expectation operator  $\mathbb{E}\left[\cdot\right]$  is a 1001-dimensional integral.
  - 1000 idiosyncratic shocks + 1 aggregate shock.
- 3. What about the stationarity of the solution and the transversality condition?
  - I will come back to this.

## General class of problems: A "big X, little x" dynamic programming

$$v(x,X) = \max_{u} \left\{ r(x,u,X) + \beta \mathbb{E} \left[ v(x',X') \right] \right\}$$
s.t.  $x' = g(x,u) + \sigma w + \eta \omega$ 

$$X' = G(X) + \Omega W + \eta \omega \mathbf{1}_{N}$$

- 1. x is the individual state of the agent.
- 2. X is a vector stacking the individual states of all of the N agents in the economy.
- 3. u is the control variable.
- 4. w is random innovation to the individual state, stacked in  $W \sim \mathcal{N}(\mathbf{0}_N, \mathbf{I}_N)$  and where, w.l.o.g.,  $w = W_1$ .
- 5.  $\omega \sim \mathcal{N}(0,1)$  is a random aggregate innovation to all the individual states.

## **Permutation Groups**

- A permutation matrix is a square matrix with a single 1 in each row and column and zeros everywhere else.
- Let  $S_N$  be the set of all n! permutation matrices of size  $N \times N$ . For example:

$$S_2 = \left\{ \begin{bmatrix} 1 & 0 \\ 0 & 1 \end{bmatrix}, \begin{bmatrix} 0 & 1 \\ 1 & 0 \end{bmatrix} \right\}$$

- Multiplying vector  $v \in \mathbb{R}^N$  by  $\pi \in S_N$  reorders elements of v
- (If you know about this):  $S_N$  is the *symmetric group* under matrix multiplication.

## Permutation-invariant dynamic programming

#### **Definition**

A 'big X, little x' dynamic programming problem is a permutation-invariant dynamic programming problem if, for all  $(x, X) \in \mathbb{R}^{N+1}$  and all permutations  $\pi \in \mathcal{S}_N$ 

1. The reward function *r* is **permutation invariant**:

$$r(x, u, \pi X) = r(x, u, X)$$

2. The deterministic component of the law of motion for X is permutation equivariant:

$$G(\pi X) = \pi G(X)$$

3. The covariance matrix of the idiosyncratic shocks satisfies

$$\pi\Omega=\Omega\pi$$

## Permutation invariance of the optimal solution

#### **Proposition**

The optimal solution of a permutation-invariant dynamic programming problem is permutation invariant. That is, for all  $\pi \in \mathcal{S}_N$ :

$$u(x, \pi X) = u(x, X)$$

and:

$$v(x, \pi X) = v(x, X)$$

Can u(x, X) permutation invariance guide  $\mathcal{H}(\Theta)$  choice?

## Representation of permutation-invariant functions

#### **Proposition**

(based on Wagstaff et al., 2019) Let  $f: \mathbb{R}^{N+1} \to \mathbb{R}$  be a continuous permutation-invariant function under  $S_N$ , i.e., for all  $(x, X) \in \mathbb{R}^{N+1}$  and all  $\pi \in S_N$ :

$$f(x,\pi X)=f(x,X)$$

Then, there exist a latent dimension  $L \leq N$  and continuous functions  $\rho : \mathbb{R}^{L+1} \to \mathbb{R}$  and  $\phi : \mathbb{R} \to \mathbb{R}^L$  such that:

$$f(x,X) = \rho\left(x, \frac{1}{N}\sum_{i=1}^{N}\phi(X_i)\right)$$

## Representation of permutation-invariant functions: Discussion and intuition

$$u(x,X) = \rho\left(x, \frac{1}{N}\sum_{i=1}^{N}\phi(X_i)\right)$$

- This proposition should remind you of Krusell-Smith (1998), L=1,  $\phi(X_i)=X_i$ .
- Key benefit for approximation is the **representation**  $(\rho, \phi)$ , **not explicit** dimensionality reduction.
- Fitting a  $\rho$  and  $\phi$  rather than f directly leads to far better generalization on  $\mathcal{X}$ . Why?:
  - Imposing structure on  $\mathcal{H}(\Theta)$ , functions that know a lot about the economic problem.
- In practice:  $L \ll N$  generalizes very well.

## **High-dimensional expectation**

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## **Expected gradient bounded in N**

#### Definition (Expected gradient bounded in N)

Let  $f: \mathbb{R}^N \to \mathbb{R}$  be a bounded function in N and  $z \sim \mathcal{N}(\mathbf{0}_N, \mathbf{I}_N)$  be a normalized Gaussian random vector. The function f has its expected gradient bounded in N if there exists a C such that:

$$\mathbb{E}\left[\|\nabla f(z)\|^2\right] \leq \frac{C}{N},$$

where C does not depend on N.

$$\mathbb{E}_{W}\left[\|\nabla u(x',X')\|^{2}\right] \leq \frac{C}{N}$$

- The policy to be well-behaved (non-explosive gradients).
- Other agent's influence vanishes.

#### Main result II: Concentration of measure

#### **Proposition**

Suppose  $z \sim \mathcal{N}(\mathbf{0}_N, \Sigma)$ , where the spectral radius of  $\Sigma$ , denoted by  $\rho(\Sigma)$ , is independent of N,  $z^1$  a draw from z, and  $f: \mathbb{R}^N \to \mathbb{R}$  is a function with expected gradient bounded in N. Then:

$$\mathbb{P}\left(\left|f(z^1) - \mathbb{E}\left[f(z)\right]\right| \ge \epsilon\right) \le \frac{\rho(\Sigma)C}{\epsilon^2} \frac{1}{N}$$

- As Ledoux (2001) puts it: "A random variable that depends in a Lipschitz way on many independent variables (but not too much on any of them) is essentially constant."
- With concentration of measure, dimensionality is not a curse; it is a blessing.

**Implication**: We can calculate  $\mathbb{E}_{W}[u(x',X')|w,\omega]$  with a *single draw* of idiosyncratic shocks W:

- $\mathbb{E}_W[u(x',X')|w,\omega] \approx u(x',X')|w,\omega$ .
- Reducing an N + 1-dimensional conditional expectation to a 2-D one (with good approximation).

## **Summarizing results**

- The structure symmetry imposes on the functions leads to better **generalization** 
  - Functions extrapolate better outside of the grid points  $\hat{\mathcal{X}}$ .
- Concentration of measures provides a fast method for calculating the conditional expectations.
  - Calculate with one draw of the idiosyncratic shocks (conditional on the aggregate shock).
- No non-heuristic algorithm exists to solve this problem.

## Solving the Model

## Design of $\mathcal{H}(\Theta)$ : Deep learning architectures

$$u(x,X) = \rho\left(x, \frac{1}{N}\sum_{i=1}^{N}\phi(X_i)\right)$$

Three cases for  $\phi$ :

- 1. Identity function: One moment  $\rightarrow \phi$  (Identity).
- 2. Up to degree four polynomials: 4 moments  $\rightarrow \phi(Moments)$ .
- 3. A deep neural network for  $\phi$ , with  $L = 4 \rightarrow \phi(\text{ReLU})$ .

If polynomials for  $\phi$ : A finite set of moments à la Krusell-Smith.

- In all cases,  $\rho$  is a highly over-parameterized neural network with four layers.
  - The baseline  $\phi$ (Identity),  $\phi$ (Moments), and  $\phi$ (ReLU) have 49.4K, 49.8K, and 66.8K coefficients.

## Solution method follows "interpolation" methods

- 1. Pick:  $\hat{X}$  as simulated trajectories from  $X_0$ :
  - Only need 100 to 1000 points regardless of dimensionality of the state space N.
  - Because we use economic insight, i.e., symmetry which gives us good generalization.
- 2. **Choose**: Design the  $\mathcal{H}(\Theta)$  with  $\rho$  and  $\phi$  as discussed:
  - $\phi$ (Identity),  $\phi$ (Moments), and  $\phi$ (ReLU).

Utilizing concentration of measures:

• One draw  $\hat{W} = \{\hat{W}_1, \dots, \hat{W}_N\}$  of the idiosyncratic shocks. For a given  $u(\cdot; \theta)$ , and aggregate shock  $\omega$  calculate:

$$X'_{i} = (1 - \delta)X_{i} + u(X) + \sigma \hat{W}_{i} + \eta \omega, \text{ for } i \in \{1, ..., N\}.$$

## Solution method follows "interpolation" methods

Approximate the Euler residuals

$$\varepsilon(X; u(\cdot; \theta)) \equiv \gamma u(X; \theta) - \beta \mathbb{E} \left[ P(X') + \gamma (1 - \delta) u(X'; \theta) \right]$$

using concentration of measures (one draw of W in X').  $\bigcirc$  error analysis in N

3. Fit: The residuals  $\varepsilon(X; u(\cdot; \theta))$ , that is the "model" i.e.,  $\ell$ .

$$\min_{\theta \in \Theta} \sum_{X \in \hat{\mathcal{X}}} \varepsilon (X; \hat{u}(\cdot; \theta))^{2}$$

4. How to Verify/Test: Given the approximate solution simulate new paths from  $X_0$  and check the Euler residuals  $(\varepsilon)$ .

Study two cases: linear (
u=1) and nonlinear (
u>1) demand functions

## Case 1: Linear to verify algorithms and methods

- With  $\nu = 1$ , we have a linear demand function:  $p(X) = 1 \frac{1}{N} \sum_{i=1}^{N} X_i$ .
- It generates a Linear-Quadratic (LQ) dynamic programming problem (only the mean of  $X_i$  matters).
- We can find the exact u(x, X), LQ has algebraic solutions.
- The LQ solution gives us a benchmark against which we can compare our deep learning solution.
- The neural network figures out very quickly that the solution is  $u(x, X) = H_0 + \frac{1}{N} H_1 \sum_{i=1}^{N} X_i$  and finds a high-dimensional approximation which matches that for the training grid.

#### **Euler residuals: Linear case**

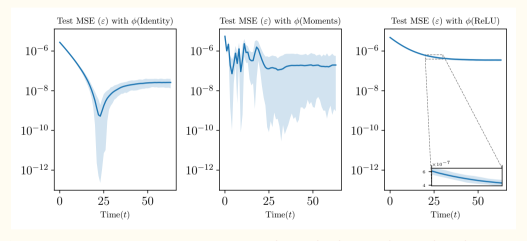


Figure 1: The Euler residuals for  $\nu=1$  and N=128 for  $\phi(\text{Identity})$ ,  $\phi(\text{Moments})$ , and  $\phi(\text{ReLU})$ . The dark blue curve shows the average residuals along equilibrium paths for 256 different trajectories. The shaded areas depict the 2.5th and 97.5th percentiles.

## **Equilibrium Paths: Linear case**

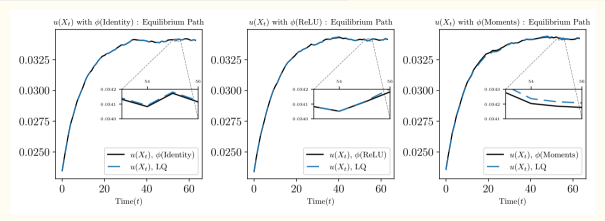
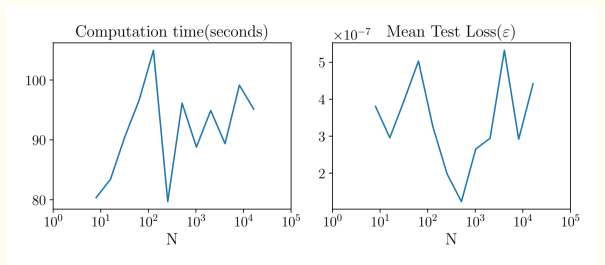


Figure 2: Comparison between baseline approximate solutions and the LQ solution for the case with  $\nu=1$  and N=128.

## Computation time: Linear case



**Figure 3:** Performance of the  $\phi(\text{ReLU})$  for different N (median value of 21 trials).

#### Case 2: Nonlinear case with no "closed-form" solution

- With  $\nu > 1$ , we have a nonlinear demand function:  $p(X) = 1 \frac{1}{N} \sum_{i=1}^{N} X_i^{\nu}$ .
- Notice how, now, the whole distribution of  $X_i$  matters.
- But we can still find the solution to this nonlinear case using exactly the same functional approximation and algorithm as before.
- ullet We do not need change anything in the code except the value of u.
- Since the LQ solution no longer holds, we do not have an exact solution to use as a benchmark, but can check residuals.
- ullet Same model and method. Computation time by  ${\it N}$  nearly the same to linear case

#### **Euler residuals: Nonlinear case**

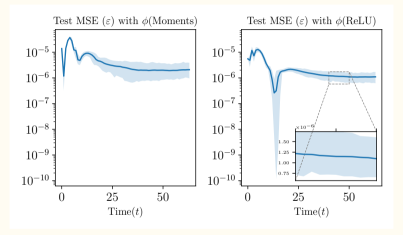
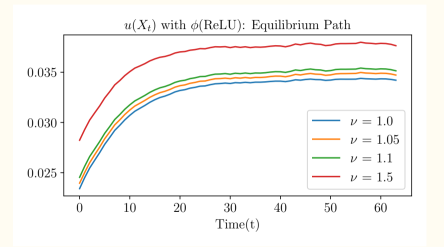


Figure 4: The Euler residuals for  $\nu=1.5$  and N=128 for  $\phi(\text{Moments})$  and  $\phi(\text{ReLU})$ . The dark blue curve shows the average residuals along equilibrium paths for 256 different trajectories. The shaded areas depict the 2.5th and 97.5th percentiles.

# Equilibrium paths: Nonlinear case



**Figure 5:** The optimal policy u along the equilibrium paths for v = [1.0, 1.05, 1.1, 1.5] and N = 128. Each path shows the optimal policy for a single trajectory.

# Some challenging question: Generalization puzzle

#### Question I: Generalization

- From statistical learning and numerical analysis we know:
  - More coefficients in the family of parametric functions  $\mathcal{H}(\Theta)$  leads to over-fitting and poor generalization (bias-variance trade-off).
  - We have 70K parameters, and < 1K grid points.
  - ullet The results indicate the opposite: More coefficients ullet better generalization.

How come we achieve great generalization?

# Some challenging questions: Multiplicity and transversality puzzle

#### Question II: Multiplicity and transversality

$$\gamma u(X) = \beta \mathbb{E} \left[ p(X') + \gamma (1 - \delta) u(X') \right]$$

$$X'_i = (1 - \delta) X_i + u(X) + \sigma W_i + \eta \omega, \quad \text{for } i \in \{1, ..., N\}$$

with linear prices. Guess and verify with  $u(X) \equiv H_0 + \frac{1}{N}H_1\sum_{i=1}^{N}X_i$ 

- The Euler equation is quadratic  $\rightarrow$  **two** solutions:  $(H_0^-, H_1^-), (H_0^+, H_1^+)$ :
  - $H_1^- < 0 o$  stationary solution,  $H_1^+ > 0 o$  non-stationary solution.
  - We have no explicit device in our algorithm to weed out the second solution.

How come we never observe the non-stationary solution in the results?

Understanding the **implicit bias** of deep neural networks answers both questions. The **Ebrahimi** Kahou et al. (2022) addresses these two challenging questions.

Implicit bias, Generalization, and

Stationarity.

### Representation with linear prices

Recall the representation,

$$u(x,X) = \rho\left(x, \frac{1}{N}\sum_{i=1}^{N}\phi(X_i)\right).$$

For  $\nu=1$  we can show that the following exact solution holds with our representation

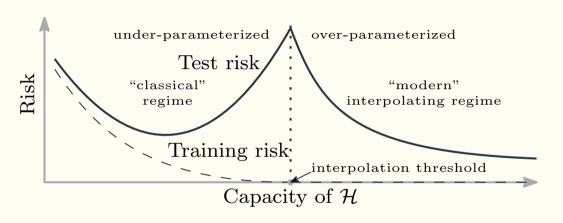
- $\phi(X_n) = X_n$  identity, L = 1.
- Doesn't matter how to generate X since only need 2 points.
- Let's do it with 3 points.

## Extreme example of generalizability of neural networks

- Extreme case : Forget we know any closed form, and see if over-fitting hurts us.
- Fit three grid points in  $\mathbb{R}^{512}$  (an economy with N = 512 agents).
- Flexible functional form with 17.7 K coefficients.
- Now, evaluate for a whole bunch of reasonable trajectories from the initial condition and check the policy error:
  - $5 \times 10^{-5}$  MSE of Euler, approximately 0.06% relative error of u(X).

This is related to a literature called **double descent**.

## The cure to over-fitting is to add more parameters



Belkin et al., 2019: Traditional statistics/bias-variance trade-off stop around the interpolation threshold.

# Deep learning optimizes in a space of functions

Remember

$$\min_{\theta \in \Theta} \sum_{\mathbf{x} \in \hat{\mathcal{X}}} \ell(\hat{\psi}(\cdot; \theta), \mathbf{x})^2$$

- Deep learning: number of coefficients is much larger than the number of grid points.
- Since  $M \gg D$ , it is possible for  $\hat{\psi}$  to interpolate and the objective value will be  $\approx 0$ .
- Since  $M \gg D$  there are many solutions (e.g.,  $\theta_1$  and  $\theta_2$ ),
  - Agree on the grid points:  $\hat{\psi}(x; \theta_1) \approx \hat{\psi}(x; \theta_2)$  for  $x \in \hat{\mathcal{X}}$ .
- ullet Since individual heta are irrelevant it is helpful to think of optimization directly within  ${\cal H}$

$$\min_{\hat{\psi}\in\mathcal{H}}\sum_{x\in\hat{\mathcal{X}}}\ell(\hat{\psi},x)^2$$

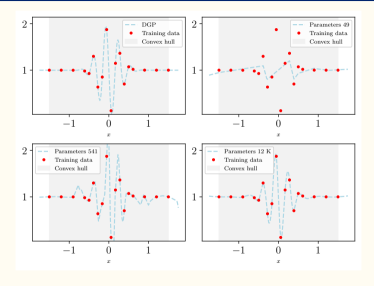
### **Deep learning and interpolation**

- For M large enough, optimizers **tend to** converge to **unique** "simple"  $\hat{\psi}$  (w.r.t to some norm  $\|\cdot\|_s$ ). Unique both in  $\hat{\mathcal{X}}$  and  $\mathcal{X}$ . There is a **bias** toward a specific class of solutions.
- How to interpret: interpolating solutions for some functional norm  $\|\cdot\|_S$

$$\min_{\hat{\psi} \in \mathcal{H}} ||\hat{\psi}||_{S}$$
 s.t.  $\ell(\hat{\psi}, x) = 0$ , for  $x \in \hat{\mathcal{X}}$ 

- Comp Sci literature refers to this as the inductive bias or implicit bias: optimization process is biased toward particular  $\hat{\psi}$ .
- Small values of  $\|\cdot\|_S$  corresponds to flat solutions with small gradients (w.r.t. input).

# Flat and smooth interpolation: Illustration



### Implicit bias: More details

Let  $\psi_1$  and  $\psi_2$  be two differentiable function from a compact space  $\mathcal X$  in  $\mathbb R$  to  $\mathbb R$  such that

$$\int_{\mathcal{X}} \left| \frac{d\psi_1}{dx} \right|^2 dx > \int_{\mathcal{X}} \left| \frac{d\psi_2}{dx} \right|^2 dx$$

then

$$\|\psi_1\|_{\mathcal{S}} > \|\psi_2\|_{\mathcal{S}}.$$

Recently shown the optimizers (first order e.g. SGD) regularize Sobolev semi-norms: Ma, Ying (2021).

### Answering the challenging questions

- Answering **generalization puzzle**: Flat interpolation leads to good generalization:
  - If the true underlying functions is flat between (and outside) the points.
  - The cure to over-fitting is to add more parameters.
- Answering multiplicity puzzle: In the linear set-up, the explosive solution has larger derivatives (less flat) than the non-explosive one i.e,  $|H_1^+| > |H_1^-|$ :
  - The deep-learning based solution automatically satisfies stationarity.
- Ebrahimi Kahou et al. (2022) explore this for many more dynamic models in macroeconomics (e.g., neoclassical growth and asset pricing) we show:
  - We can have short- and medium-run accurate solutions without being worried about the long-run behavior.
  - We dont need to calculate the steady-state (ergodic distribution).

# Conclusions

#### Extensions

- 1. Decreasing returns to scale: the policy becomes a function of x.
- 2. Multiple productivity types (e.g., two different groups).
- 3. Complex idiosyncratic states (e.g., a agent is described with more than one variable).

#### **Summarizing our contribution**

- **Method** for solving **high-dimensional** dynamic programming problems and competitive equilibria with idiosyncratic and aggregate shocks relying
  - Symmetry.
  - Concentration of measures: Dimensionality is a blessing not a curse.
- Using economic theory (i.e., exchangeability) and deep learning for function approximation with a huge # of parameters (>> grid points)
  - Achieve great generalization: key to alleviate the curse of dimensionality.
- Implementation
  - Can deal with 10000+ agents.
  - Can deal with 10000+ dimensional expectations with one Monte-carlo draw.

# **Appendix**

#### Definition (Bounded functions in N)

Let:

$$\mathcal{L}(M) \equiv \{ y \in \mathbb{R}^N : |y_i| \le M \ \forall i = 1, \dots, N \}$$

be an N-dimensional hypercube in  $\mathbb{R}^N$ . A function  $f: \mathbb{R}^N \to \mathbb{R}$  is bounded in N if for every M there exists  $K_M$  such that

$$\sup_{y \in \mathcal{L}(M)} |f(y)| < K_M,$$

where  $K_M$  is a constant that does not depend on N, but may depend on M.

- Example  $f(y) = \frac{1}{N} \sum_{i=1}^{N} y_i \rightarrow \sup_{y \in \mathcal{L}(M)} |f(y)| < M$ .
- To avoid  $f(y) = \sum_{i=1}^{N} y_i \to \sup_{y \in \mathcal{L}(M)} |f(y)| < NM$ .



## Concentration of measure is the bless of dimensionality

In the linear case we know the closed form solution for u

$$\hat{\varepsilon}(X; u) - 0 \sim \mathcal{N}\left(0, \frac{\sigma_{\varepsilon}^{2}}{N}\right)$$
$$u(\hat{X}') - \mathbb{E}\left[u(X') \mid \omega\right] \sim \mathcal{N}\left(0, \frac{\sigma_{u}^{2}}{N}\right)$$

- Conditional expectation becomes constant as N gets large.
  - One single Monte-carlo draw of the idiosyncratic shocks is enough.



#### Analytic euler error due to the concentration of measure

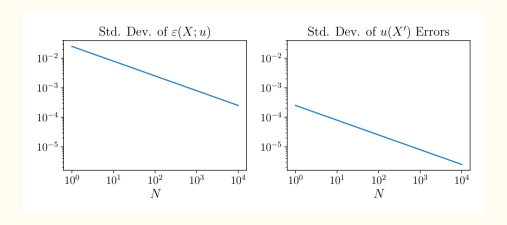


 Table 1: Performance of Different Networks in Solving the Linear Model

		Time (s)	Params (K)	Train MSE $(\varepsilon)$	Test MSE $(\varepsilon)$	Val MSE $(\varepsilon)$	Policy Error $( u - u_{ref} )$	Policy Error $\left(\frac{ u-u_{\text{ref}} }{u_{\text{ref}}}\right)$
group	description							
$\phi$ (Identity)	Baseline	42	49.4	4.1e-06	3.3e-07	3.3e-07	2.9e-05	0.10%
	Thin (64 nodes)	33	12.4	3.7e-06	2.7e-07	2.7e-07	3.4e-05	0.10%
$\phi(Moments)$	Baseline	55	49.8	1.4e-06	7.6e-07	7.6e-07	2.8e-05	0.09%
	Moments (1,2)	211	49.5	2.4e-06	1.1e-06	2.3e-06	4.4e-05	0.14%
	Very Shallow(1 layer)	241	0.6	1.1e-05	8.4e-06	7.9e-06	1.1e-02	34.00%
	Thin (64 nodes)	82	12.6	1.6e-06	9.1e-07	9.2e-07	3.8e-05	0.12%
	Baseline	107	66.8	3.7e-06	3.3e-07	3.3e-07	2.7e-05	0.09%
$\phi(ReLU)$	L = 2	86	66.3	1.3e-05	2.1e-07	2.2e-07	2.6e-05	0.08%
	L = 16	91	69.9	5.5e-06	1.5e-07	1.5e-07	2.1e-05	0.07%
	Shallow ( $\phi$ : 1 layer, $\rho$ : 2 layers)	79	17.7	2.0e-06	5.5e-07	5.5e-07	3.2e-05	0.11%
	Thin $(\phi, \rho$ : 64 nodes)	87	17.0	1.1e-05	4.5e-07	4.5e-07	3.0e-05	0.10%

Table 2: Nonlinear Model Performance

		Time (s)	Params (K)	Train MSE $(\varepsilon)$	Test MSE $(\varepsilon)$	Val MSE $(\varepsilon)$
group	description					
	Baseline	26	49.8	6.0e-06	5.0e-06	3.8e-06
	Moments (1,2)	27	49.5	8.0e-06	5.1e-06	3.6e-06
4(Momenta)	Very Shallow (1 layer)	252	0.6	8.3e-06	1.4e + 00	5.0e-06
$\phi$ (Moments)	Thin (32 nodes)	66	3.2	1.1e-05	9.7e-06	4.4e-06
	Baseline	60	67.1	1.4e-05	4.7e-06	3.3e-06
	L = 8	73	68.1	1.1e-05	4.9e-06	2.0e-06
	L = 16	72	70.2	1.5e-05	5.4e-06	1.7e-06
$\phi(ReLU)$	Very Shallow $(\phi,  ho: 1$ layer)	136	1.4	8.9e-06	4.8e + 06	4.9e-06
$\psi(\text{ReLO})$	Shallow $(\phi, \rho: 2 \text{ layers})$	47	34.3	1.0e-05	9.2e-06	2.8e-06
	Thin $(\phi, \rho : 32 \text{ nodes})$	52	4.5	1.3e-05	6.0e-06	2.7e-06

#### **Parameters**

• 
$$\gamma=$$
 90,  $\beta=$  0.95,  $\sigma=$  0.005,  $\eta=$  0.001.