

Deep Learning Solutions to Master Equations for Continuous Time Heterogeneous Agent Macroeconomic Models

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March 21, 2023

Abstract

We propose new global solution algorithms for continuous time heterogeneous agent economies with aggregate shocks. We first approximate the state space so the master equation becomes a high, but finite, dimensional partial differential equation. We then approximate the value function using neural networks and solve the master equation using deep learning tools. The main advantage of this technique is that it allows us to find global solutions to high dimensional, non-linear problems. We demonstrate our algorithms by solving two canonical models in the macroeconomics literature: the [Aiyagari \(1994\)](#) model and the [Krusell and Smith \(1998\)](#) model.

Keywords: Heterogeneous agents, computational methods, deep learning, inequality, mean field games, continuous time methods, aggregate shocks, global solution.

1 Introduction

There is great interest in understanding how inequality changes over time and reacts to government policy. This has led researchers to integrate the evolution of population distributions into macroeconomic models. One major difficulty with working on these models is that the agent distribution becomes a state variable and so the state space becomes infinite dimensional. In this paper, we demonstrate how deep learning techniques can relax the “curse of dimensionality” for continuous time heterogeneous agent models and allow global numerical solutions to be computed.

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We develop solution techniques for a class of dynamic, stochastic, general equilibrium economic models with the following features. There is a large collection of price-taking agents who face both idiosyncratic and aggregate shocks. Given their belief about the evolution of aggregate state variables, agents choose control processes to solve dynamic optimization problems. When making their decisions, agents face financial frictions that constrain their behaviour and potentially break “aggregation” results that would allow the distribution of agents to be replaced by a “representative agent”. We solve for a rational expectations equilibrium in which agent beliefs about the evolution of aggregate states are consistent with the dynamics that emerge in the economy. Solving for equilibrium reduces to solving a “master” partial differential equation (pde) that summarizes both agent optimization behaviour (from the Hamilton-Jacobi-Bellman equation) and the evolution of the distribution (from the Kolmogorov forward equation). A canonical example of this type of environment is the [Krusell and Smith \(1998\)](#) model, which is often used as a workhorse environment for testing solution methods in macroeconomics.

Our solution approach approximates the infinite dimensional master equation by a finite, but high, dimensional pde and then uses deep learning to solve the high dimensional equation. We consider two different approaches for reducing the dimension of the master equation. The first approach approximates the distribution by a large, finite number of agents. We refer to this as the “finite-agent” approximation. The second approach approximates the distribution by discretizing the agent state space so the density becomes a collection of mass points at grid points. We call this the “finite-state” approximation.

We solve the finite dimensional approximation to the master equation using recent advances in deep learning. In particular, we adapt the Deep Galerkin Method (DGM) from the applied mathematics literature. This approach approximates the value function by a neural network and then uses stochastic gradient descent to train the neural network to minimise a loss function that combines the average error in master equation with the average error in the boundary conditions. We calculate average errors by randomly sampling over points in the state space and the boundary, with greater sampling applied to the regions with more curvature. We also need to handle inequality boundary constraints arising from financial constraints on the evolution of the state space. We do this by introducing penalty functions for when financial constraints bind.

We illustrate our technique by solving two canonical models in the macroeconomics literature that are frequently used for testing numerical techniques: [Aiyagari \(1994\)](#) and [Krusell and Smith \(1998\)](#). For the [Aiyagari \(1994\)](#) model, we show that we can match the finite difference solution to high accuracy and solve for transition dynamics without a shooting algorithm. For the [Krusell and Smith \(1998\)](#) model, we show that we generate solutions with a low error for the master equation. Traditional techniques only give approximate solutions to the [Krusell and Smith \(1998\)](#) model so we do not have a ideal benchmark for testing the model. However, we show that we get very similar results to contemporary approaches such as [Fernández-Villaverde et al. \(2018\)](#).

Literature Review: The economics literature has traditionally used three main approaches for solving heterogeneous agent models with aggregate shocks. One approach is to fit a statistical approximation to the law of motion for the key aggregate state variables (e.g. [Krusell and Smith \(1998\)](#), [Den Haan \(1997\)](#), [Fernández-Villaverde et al. \(2018\)](#)). As has been extensively discussed in the literature, this approach works well when the law of motion for the key state variables can be well approximated as a function of key moments of the distribution (and so the economy is very close to permitting “aggregation”). By contrast, our approach can handle economies without near aggregation results. A second approach is to take a type of linear perturbation in the aggregate state and then solve the resulting linear problem with matrix algebra (e.g. [Reiter \(2002\)](#), [Reiter \(2008\)](#), [Reiter \(2010\)](#), [Winberry \(2018\)](#), [Ahn et al. \(2018\)](#), [Auclert et al. \(2021\)](#), [Bilal \(2021\)](#), [Bhandari et al. \(2023\)](#)). By contrast, we solve the model globally and so can handle partial differential equations with extensive non-linearity. A final approach is to take a low dimensional projection of the distribution (e.g. [Prohl \(2017\)](#), [Schaab \(2020\)](#)). Our approach is complementary to these papers in that it allows for more general, higher dimensional projections.

Our paper is part of a growing computational economics literature using deep learning techniques to solve economics models and overcome the difficulties of the traditional solution techniques. Many of these papers focus on solving heterogeneous agent macroeconomic models in discrete time (e.g. [Azinovic et al. \(2022\)](#), [Han et al. \(2021\)](#), [Maliar et al. \(2021\)](#), [Kahou et al. \(2021\)](#), [Bretscher et al. \(2022\)](#)). We are part of a less developed literature attempting to deploy deep learning techniques to continuous time models (e.g. [Duarte \(2018\)](#), [Gopalakrishna \(2021\)](#), [Fernandez-Villaverde et al. \(2020\)](#)). Our contribution to the continuous time literature is to understand how to handle a rich distribution of agents and directly solve the “master equation” for the economic system. This necessitates resolving a collection of problems that are particular to continuous time such as: approximating inequality boundary conditions, using the Kolmogorov Forward Equation to derive laws of motion for projection coefficients, and approximating derivatives with respect to the distribution.

There is also a growing mathematics literature that attempts to use neural networks to solve differential equations. Technically, our approach builds on the Deep Galerkin Method developed in [Sirignano and Spiliopoulos \(2018\)](#) and [Li et al. \(2022\)](#). Our approach to handling borrowing constraints draws on [Lu et al. \(2021b\)](#) and [Brzoza-Brzezina et al. \(2015\)](#). We also build on the literature using neural networks to solve mean field games. [Al-Arabi et al. \(2022\)](#); [Carmona and Laurière \(2021\)](#) adapted the Deep Galerkin Method to solve the PDE system arising in mean field games. [Fouque and Zhang \(2020\)](#); [Carmona and Laurière \(2022a\)](#); [Germain et al. \(2022b\)](#) proposed deep learning methods for mean field games and mean field control problems based either on direct approximation of the control or on an adaptation of the Deep BSDE method proposed by [Han et al. \(2018\)](#). We refer to e.g. [Laurière \(2021\)](#); [Carmona and Laurière \(2022b\)](#); [Hu and Lauriere \(2022\)](#) and the

references therein for more details. However, these works are mostly focused on solving the problem at equilibrium and without aggregate shocks. [Min and Hu \(2021\)](#) proposed a deep learning method based on recurrent neural networks and signatures to solve mean field games with aggregate shocks, when the interactions are through moments and without solving the Master equation. [Perrin et al. \(2022\)](#) introduced a deep reinforcement learning algorithm based on fictitious play to learn population-dependent policies for finite-state, finite-action mean field games. In the context of mean field control, [Carmona et al. \(2019\)](#); [Gu et al. \(2021\)](#); [Germain et al. \(2022a\)](#) used deep learning to compute social optima but these methods do not solve Nash equilibria. Relative the mathematics literature, our focus is on solving Master equations for a class of mean field games with aggregate shocks.

This document is organised as follows. Section 2 describes the general economic environment that we will be studying and derives the master equation. Section 3 describes the different approaches to solving the model. Section 4 applies our algorithm to the continuous time version of the [Aiyagari \(1994\)](#) described in [Achdou et al. \(2022\)](#). Section 5 applies our algorithm to continuous time version of [Krusell and Smith \(1998\)](#). Section 6 dicusses practical lessons. Section 7 concludes.

2 Economic Model

In this section, we outline the class of economic models for which our techniques are appropriate. At the high level, in economics terminology, we are solving continuous time, general equilibrium models with a distribution of optimizing agents who face idiosyncratic and aggregate shocks.¹ In mathematics terminology, we are solving mean field games with common noise.

Setting: The model is in continuous time with infinite horizon. There is an exogenous one-dimensional aggregate state variable, $z_t \in \mathcal{Z}$, which evolves according to:

$$dz_t = \mu^z(z_t)dt + \sigma^z(z_t)dB_t^0, \quad (2.1)$$

where B_t^0 denotes a common Brownian motion process.

Agent Problem: The economy is populated by a collection of agents, denoted by I . Each agent, $i \in I$, has an idiosyncratic state vector, $x^i \in \mathcal{X}$, that evolves according to:

$$dx_t^i = \mu^x(c_t^i, x_t^i, z_t, q_t)dt + \sigma^x(x_t^i, z_t, q_t)dB_t^i + \gamma^x(x_t^i, z_t, q_t)dJ_t^i, \quad (2.2)$$

where $c_t^i \in \mathcal{C}$ is a one-dimensional control variable chosen by the agent, $u(c_t^i)$ is the flow utility from choosing c_t^i , $q_t \in \mathcal{Q}$ is a collection of aggregate prices in the economy that will

¹For ease of exposition, restrict attention here to models with 1-dimensional aggregate shocks. We set the Master equation for the full class of models in Appendix A.1.

be determined endogeneously in equilibrium, B_t^i denotes an N -dimensional idiosyncratic Brownian motion process, and J^i denotes an idiosyncratic Poisson jump process. We let $\lambda(x)$ denote the rate at which Poisson jump shocks arrive given idiosyncratic state x . We let $G_t = \mathcal{L}(x_t^i | \mathcal{F}_t^0)$ and g_t denote the population distribution and density across x_t^i at time t , for a given history \mathcal{F}_t^0 , where \mathcal{F}_t^0 is the filtration generated by the aggregate shock process B^0 .

Each agent, i , has a belief about the stochastic price process $\hat{q} = \{\hat{q}_t : t \geq 0\}$ adapted to \mathcal{F}_t^0 . Given their belief, agent i chooses their control process, $c^i = \{c_t^i : t \geq 0\}$, to solve:

$$V(x_0^i, z_0) = \max_{c^i} \mathbb{E}_0 \left[\int_0^\infty e^{-\rho t} u(c_t^i) dt \right] \quad (2.3)$$

s.t. (2.1), (2.2), $\Phi(x_t^i) \geq 0$,

where $\Phi(x_t^i) \geq 0$ is an exogenous constraint on the state space process that restricts the agent's choice of control process. A classic example in economics problems is the constraint that x_t^i is bounded below: $\Phi(x_t^i) = x_t^i - \underline{x} \geq 0$. We let \mathcal{X} denote the domain of x_t^i implicitly defined by the constraint that $\Phi(x_t^i) \geq 0$.

Equilibrium: Given an initial density g_0 , an equilibrium for this economy consists of a collection of \mathcal{F}_t^0 -adapted stochastic process, $\{c_t^i, g_t, q_t, z_t : t \geq 0, i \in I\}$, that satisfy the following conditions: (i) each agent's control process c_t^i solves problem (2.3) given their belief that the price process is \hat{q} , (ii) the equilibrium prices satisfy a market clearing condition that connects the prices to the distribution of agents:

$$q_t = Q(z_t, \bar{g}_t), \quad \forall t \geq 0,$$

where \bar{g}_t is the mean of g , and (iii) agent beliefs about the price process are consistent with the optimal behaviour of other agents in the sense that $\hat{q} = q$.

Master Equation: We assume that a solution to the equilibrium exists that is recursive in the aggregate state variables: $\{z, g\}$. We follow the approach of [Lions \(2011\)](#) and characterize the equilibrium in one PDE, which is referred to as the “master equation” of the “mean-field-game”. This formulation is particularly convenient when the evolution of the economy is subject to aggregate shocks and the evolution of the aggregate state variables cannot be determined deterministically. Denote the recursive equilibrium optimal control of the individual agents by $c^*(x, z, g)$. Then, for a given z path, the evolution of the distribution under the optimal control $c^*(x, z, g)$ be characterized by the Kolmogorov Forward Equation

(KFE) of the form²:

$$dg_t(x) = \mu_g(c^*(x, z, g), z_t, g_t)dt. \quad (2.4)$$

Let $V(x, z, g)$ denote the solution for the following PDE on the space $x \in \mathcal{X}$:

$$(\mathcal{L}V)(x, z, g) = 0 \quad (2.5)$$

where the operator $(\mathcal{L}V)(x, z, g) := (\mathcal{L}^hV + \mathcal{L}^gV)(x, z, g)$ is defined by:

$$\begin{aligned} (\mathcal{L}^hV)(x, z, g) := & -\rho V(x, z, g) + u(c^*(x, z, g)) \\ & + D_xV(x, z, g)\mu^x(c^*(x, z, g), x, z, Q(z, \bar{g})) + \partial_zV(x, z, g)\mu^z(z) \\ & + \frac{1}{2}\text{tr}\{\Sigma^x(x, z, Q(z, \bar{g}))D_x^2V(x, z, g)\} + \frac{1}{2}(\sigma^z(z))^2\partial_{zz}V(x, z, g) \\ & + \lambda(x)(V(x + \gamma^x(x, z, g), z, g) - V(x, z, g)) \\ (\mathcal{L}^gV)(x, z, g) := & \int_{\mathcal{X}} \mu_g(c^*(x, z, g), z, g) \frac{\partial V}{\partial g}(x, z, g)(y)dy. \end{aligned}$$

where \mathcal{L}^h reflects the optimization problem of the household, \mathcal{L}^g reflects how the evolution of the distribution affects the household value, $\Sigma^x(\cdot) := \sigma^x(\cdot)(\sigma^x(\cdot))^T$, $\partial V/\partial g$ is the Frechet derivative of V with respect to the distribution, and the master equation is subject to a problem-specific boundary condition related to the constraint $\Phi(x_t^i) \geq 0$.³

Intuitively, $V(x, z, g)$ can be interpreted as the optimal value of a representative player who starts at state x , with aggregate shock equal to z , and who faces a population that starts at the distribution g and then plays according to the Nash equilibrium control c . We refer to [Cardaliaguet et al. \(2015\)](#); [Bensoussan et al. \(2015\)](#) for more details. From V , the optimal consumption c^* can be computed for every (x, z, g) , which allows a representative player to react optimally to any population distribution. The equilibrium control, c^* , is characterised by the equation:⁴

$$0 = u'(c^*(x, z, g)) + D_xV(x, z, g)\partial_c\mu^x(c^*(x, z, g), x, z, Q(z, \bar{g})).$$

The goal of this paper is use deep learning techniques to find numerical solutions to equation (2.5). The challenge is that the master equation contains an infinite dimensional derivative with respect to the distribution g . We need to work with numerical approximations of the distribution and solution methods that can handle high dimensions.

²Observe that there is no noise in the KFE because dB_t^0 does not directly impact the evolution of idiosyncratic states. We consider this case for notational simplicity. The techniques outlined in this paper still apply if we relax this assumption.

³We define the gradient $D_xV(\dots)$ as a row vector, such that the product $D_xV(\dots)\mu^x(\dots)$ has to be interpreted as an inner product for multi-dimensional x .

⁴This equation can be derived from the first-order condition for c in the agent's HJB equation.

3 Generic Solution Approach

In this section, we outline a generic approach for applying a “Deep Galerkin” approach to solving the master equation (2.5). The first part of this approach is to find a finite dimensional approximation to the distribution so we can develop a finite, but high, dimensional approximation to the master equation. The second part is to approximate the solution to the finite dimensional master equation using a neural network. Finally, we use “deep-learning” to solve the approximate master equation.

3.1 Finite Dimensional Master Equation

In order to proceed, we need to find a finite dimensional approximation to the density, which we denote by \hat{g} , and the associated finite dimensional master equation operator, $\hat{\mathcal{L}}$. In this section, we outline two possible approximation approaches. The first approach approximates the distribution with a finite collection of agents. The second approach projects the distribution onto a collection of mass points on a discretized grid. The difference between the approaches primarily appears in how the operator \mathcal{L}^g is approximated.

3.1.1 Finite Agent Approximation

In this approach, we restrict the general model so that there are $I < \infty$ agents, denoted by $i \in \{1, 2, \dots, I\}$. Under the finite agent approximation, the density, g_t , is replaced by the individual states of the I agents:

$$\hat{g}_t := \{x_t^i : i \leq I\}$$

and so the clearing conditions become:

$$q_t = Q(z_t, \bar{\hat{g}}_t).$$

where $\bar{\hat{g}}_t = \frac{1}{I} \sum_{i=1}^I x_t^i$ is the average position of the agents.

To maintain the price taking assumption in the infinite agent model, we impose that agent i behaves as if their individual actions do not influence prices. Formally, this means their belief about the pricing function will be given by:

$$q_t = Q(z_t, \bar{\hat{g}}_t^{-i})$$

with $\bar{\hat{g}}_t^{-i} = \{x_t^j \in I^{-i}\}$ and $\bar{\hat{g}}_t^{-i} = \frac{1}{I-1} \sum_{j \in I^{-i}} x_t^j$, where $I^{-i} = \{j \leq I : j \neq i\}$ denotes the agents excluding agent i . Ultimately, this will ensure that the neural network trains the policies rules as if the agents believe that their assets do not influence the market prices. Aside from this change to the belief process, the optimization problem for household remains the same as in the general model.

Let $c^*(x^i, z, \hat{g})$ denote the equilibrium optimal control. Let $V(x^i, z, \hat{g})$ denote the value function for the master equation in the economy with I price taking agents. Then $V(x^i, z, \hat{g})$ solves $(\hat{\mathcal{L}}V)(x^i, z, \hat{g}) = 0$ subject to the boundary conditions, where the master equation operator is:

$$\begin{aligned} (\hat{\mathcal{L}}V) &= (\hat{\mathcal{L}}^h V) + (\hat{\mathcal{L}}^g V), \quad \text{where} \\ (\hat{\mathcal{L}}^h V)(x^i, z, \hat{g}) &:= (\mathcal{L}^h V)(x^i, z, \hat{g}) \\ (\hat{\mathcal{L}}^g V)(x^i, z, \hat{g}) &= \sum_{j \leq I} \frac{\partial V}{\partial g^j}(x^i, z, \hat{g}) \mu^x(c^*(x^j, z, \hat{g}), x^j, z, Q(z, \hat{g}^{-j})) \\ &\quad + \sum_{j \leq I} \frac{1}{2} \text{tr} \left\{ \Sigma^x(x^j, z, Q(z, \hat{g}^{-j})) D_{g^j}^2 V(x^i, z, \hat{g}) \right\} \\ &\quad + \sum_{j \leq I} \lambda(x^j) \left(V(x^i, z, \{x^j + \gamma^x(x^j, z, Q(z, \hat{g}^{-j})), \hat{g}^{-j}\}) - V(x^i, z, \hat{g}^{-i}) \right), \end{aligned}$$

subject the constraint $\Phi(x^i) \geq 0$. The operator for the household optimization problem, $\hat{\mathcal{L}}^h$, is the same as in the general problem but with the distribution replaced by the finite collection of agents for the calculation of moments to get the pricing functions. The operator for the impact of distributional changes on the household, $\hat{\mathcal{L}}^g$, changes to capture impact of changes in other agent positions.

3.1.2 Discrete State Space Approximation

In this section, we outline the second approach: projecting the distribution onto a finite collection of mass points so it be approximated by the distribution by a histogram, i.e., a vector of weights on a finite number of points. We consider N^x points, denoted by x_1, \dots, x_{N^x} , in \mathcal{X} . We will approximate g by a vector $\hat{g} \in \mathbb{R}^{N^x}$ of values at x_1, \dots, x_{N^x} . The KFE under optimal control (2.4) is replaced by an ordinary differential equation in dimension N^x of the form:

$$\dot{\hat{g}}_t = \hat{\mu}_{\hat{g}}(z_t, \hat{g}_t). \quad (3.1)$$

The right-hand-side needs to be approximated using information from the Kolmogorov Forward Equation. In our numerical examples we use a finite difference approximation to the Kolmogorov Forward Equation to derive $\hat{\mu}_{\hat{g}}$. However, the technique can be applied to other types of projections.

The solution V to the master equation is replaced by a function $V : \mathcal{X} \times \mathcal{Z} \times \mathbb{R}^{N^x} \rightarrow \mathbb{R}$

which solves the following PDE:

$$\begin{aligned}
(\hat{\mathcal{L}}V) &= (\hat{\mathcal{L}}^h V) + (\hat{\mathcal{L}}^g V), \quad \text{where} \\
(\hat{\mathcal{L}}^h V)(x, z, \hat{g}) &:= (\mathcal{L}^h V)(x, z, \hat{g}) \\
(\hat{\mathcal{L}}^g V)(x, z, \hat{g}) &= \sum_{i=1}^{N^x} [\hat{\mu}_{\hat{g}}(z, \hat{g})]_i \frac{\partial V}{\partial \hat{g}_i}(x, z, \hat{g}),
\end{aligned}$$

subject to the constraint that $\Phi(x) \geq 0$.

3.2 Neural Network Approximations

Both approaches in section 3.1 lead to finite approximations to the density, \hat{g} , and the master equation operator $\hat{\mathcal{L}}$. However, the resulting master equations are high dimensional and so cannot be solved by traditional techniques. Instead, we approximate the solution to the master equation using a neural network and deploy tools from the “deep learning” literature to “train” the neural network to solve the approximate master equation.

A neural network is a type of parametric functional approximation that is built by composing affine and non-linear functions in a chain or “network” structure (see [Goodfellow et al. \(2016\)](#) for a detailed discussion). We let $\hat{X} := \{x, z, \hat{g}\}$ denote the collection of inputs into the approximate value function. We denote the neural network approximation to the value function by $V(\hat{X}) \approx \hat{V}(\hat{X}; \theta)$, where θ are the parameters in the neural network approximation that depend upon the form of the approximation. There are many types of neural network approximations. The simplest form is a “feedforward” or “deep feedforward” neural network which is defined by:

$$\begin{aligned}
\mathbf{h}^{(1)} &= \phi^{(1)}(W^{(1)} \hat{X} + \mathbf{b}^{(1)}) && \dots \text{Hidden layer 1} \\
\mathbf{h}^{(2)} &= \phi^{(2)}(W^{(2)} \mathbf{h}^{(1)} + \mathbf{b}^{(2)}) && \dots \text{Hidden layer 2} \\
&\vdots && \\
\mathbf{h}^{(H)} &= \phi^{(H)}(W^{(H)} \mathbf{h}^{(H-1)} + \mathbf{b}^{(H)}) && \dots \text{Hidden layer H} \\
\mathbf{o} &= W^{(H+1)} \mathbf{h}^{(H)} + \mathbf{b}^{(H+1)} && \dots \text{Output layer} \\
\hat{V} &= \phi^{H+1}(\mathbf{o}) && \dots \text{Output}
\end{aligned} \tag{3.2}$$

where the $\{\mathbf{h}^{(i)}\}_{i \leq H}$ are vectors referred to as “hidden layers” in the neural network, $\{W^{(i)}\}_{i \leq (H+1)}$ are matrices referred to as the “weights” in each layer, $\{\mathbf{b}^{(i)}\}_{i \leq (H+1)}$ are vectors referred to as the “biases” in each layer, $\{\phi^{(i)}\}_{i \leq (H+1)}$ are non-linear functions applied element-wise to each affine transformation and referred to as “activation functions” for each layer. The length of hidden layer, $\mathbf{h}^{(i)}$, is referred to as the number of *neurons* in hidden layer i . The total collection of parameters is denoted by $\theta = \{W^{(i)}, \mathbf{b}^{(i)}\}_{i \leq (H+1)}$. The goal of deep learning is to train the parameters, θ , to make $\hat{V}(\hat{X}; \theta)$ a close approximation

to $V(\hat{X})$.

The neural network defined in (3.2) is called a “feedforward” because hidden layer i cannot depend on hidden layers $j > i$. This is in contrast to a “recursive” neural networks where any hidden layer can be a function of any other hidden layer. It is called “fully connected” if all the entries in the weight matrices can be non-zero so each layer can use all the entries in the previous layer. In this paper, we will consider a fully connected “feedforward” network to be the default network. This is because these networks are the quickest to train and so we typically start by trying out this approach. However, there are applications where we find that more complicated neural network formulations are useful. In particular, we find that the type of recursive neural network suggested by the “Deep Galerkin” approach in [Sirignano and Spiliopoulos \(2018\)](#) is helpful for finite state space approximations.

3.3 Solution Algorithm

We train the neural network to learn parameters θ that minimize the error in the master equation and boundary conditions. We describe the key steps in in Algorithm 1. Essentially, the algorithm generates random points in the discretized states space $\{x, z, \hat{g}\}$, then calculates the error in the master equation on those points, and updates the parameters to decrease the error in the master equation. In the deep learning literature, this approach is sometimes referred to as “unsupervised” learning (e.g. [Azinovic et al. \(2022\)](#)) because we do not have direct observations of the value function, $V(x, z, \hat{g})$, and instead have to learn the value function indirectly via the master equation.

There are some features of the algorithm that are typical to deep learning problems but less common in other economics techniques so address them here:

1. *Why do we draw a new sample each epoch rather than fixing a sample from the start?*
We find that fixing the sample across epochs leads to overfitting problems where the neural network matches the sample points well but interpolates poorly in the rest of the state space.
2. *Could the same algorithm be run with an alternative parametric approximation such as Chebyshev polynomials?* In principle, it is possible. But, a key feature of the training algorithm is that we need to be able to calculate automatic derivatives. Chebyshev polynomials have a smaller number of parameters but complicated basis functions whereas neural networks have a large number of parameters but simple basis functions. In addition, the machine literature has invested heavily in getting non-linear optimizers to work well with neural nets. For all these reasons, it is easier to run the non-linear optimizer with neural networks.
3. *Does this algorithm solve for the global or local minimum?* The stochastic gradient decent algorithm calculates the loss on random collections of points and so has

some ability to wander the parameter space looking for the global minimum. We don't always need to find the global minimum of the local minima are reasonable approximations.

Algorithm 1: Generic Solution Algorithm

1. Approximate the value function by a neural network: $V(x, z, \hat{g}) \approx \hat{V}(x, z, \hat{g}; \theta)$, where θ are the neural network parameters for the value function
2. Make initial parameter guess θ^0 .
3. At iteration n with guess θ^n :
 - (a) Generate M^e sample points, $S^{ne} = \{(x_m, z_m, \hat{g}_m)\}_{m \leq M^e}$ for evaluating the master equation error and M^b sample points, $S^{nb} = \{(\underline{x}, z_m, \hat{g}_m)\}_{m \leq M^b}$ for evaluating the boundary condition error. Let $S^n := \{S^{ne}, S^{nb}\}$ denote the collection of sample points at iteration n .

- (b) Calculate the weighted average error:

$$\mathcal{E}(\theta^n, S^n) = \kappa^e \mathcal{E}^e(\theta^n, S^{ne}) + \kappa^b \mathcal{E}^b(\theta^n, S^{nb})$$

where κ^e and κ^b are weights on the master equation error, \mathcal{E}^e , and boundary condition error, \mathcal{E}^b , and the errors are typically taken to be mean-squared errors:

$$\begin{aligned} \mathcal{E}^e(\theta^n, S^{ne}) &:= \frac{1}{M^e} \sum_{m \leq M^e} |\hat{\mathcal{L}}(x_m, z_m, \hat{g}_m)|^2 \\ \mathcal{E}^b(\theta^n, S^{nb}) &:= \frac{1}{M^b} \sum_{m \leq M^b} |\Phi(\underline{x}, z_m, \hat{g}_m)^-|^2 \end{aligned}$$

where $(\cdot)^-$ denotes the negative part and the derivatives in the differential operator and boundary condition are calculated using automatic differentiation.

- (c) Update the the parameters using “deep learning” toolkit. We typically use a “stochastic gradient descent” style method: at each point:

$$\theta^{n+1} = \theta^n - \alpha_n D_\theta \mathcal{E}(\theta^n, S^n)$$

where α_n is the “learning rate” and $D_\theta \mathcal{E}$ is the vector differential operator.

- (d) Repeat until $\mathcal{E}(\theta^n, S^n) \leq \epsilon$ where ϵ is a precision threshold.
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At a high level, the algorithm is straightforward. However, successfully implementing the deep learning approach is not trivial and involves many problem specific adjustments. In the following sections, we work through canonical macroeconomic problems in detail and compare different ways of implementing the algorithm.

3.4 Advantages of Neural Networks For Continuous Time

There are a number of advantages to using neural networks to solve differential equations, some of which are particular to continuous time problems. One advantage, as we have mentioned, is that we can deal with high dimensional differential equations. This allows us to work with high dimensional approximations to distributions that traditional techniques (e.g. finite difference and spectral methods) cannot solve in reasonable time. As we show in the following examples, this allows us to solve models with heterogeneous agents and aggregate shocks.

A second advantage is that we are able to use automatic differentiation to calculate the derivatives to evaluate the differential equations. In continuous time, the master equation contains a differential operator and so numerical techniques need to focus on efficient ways of calculating derivatives. This is very different to discrete time, where the Bellman equation has expectation operators and so numerical approaches need to focus on efficient ways of calculating expectations. Traditional continuous time techniques, especially finite difference, typically have difficulty working with regions with high curvature because the finite difference approximation to the derivative breaks down. Neural network techniques are particularly useful for continuous time because they can use automatic differentiation rather than working with discrete state space approximations to derivatives.

A third advantage is that neural networks work well with randomly sampling state space points. When using spectral methods, such as Chebyshev polynomials, the grid for the state space need to be carefully chosen to prevent synchronisation in the oscillations of the polynomials. By contrast, for neural networks, we can sample randomly from the state space and so use active learning to focus on regions with large errors.

4 Example: Uninsurable Income Risk

A canonical macroeconomic model with heterogeneous agents is the continuous time version of Aiyagari (1994) (referred as the Aiyagari-Bewley-Huggett (ABH) model in the formulation in Achdou et al. (2017) that we follow in this paper). We start by using our solution technique to solve this model because there are existing accurate solution techniques and so we can compare our solution technique in detail.

4.1 Model Specification

In this subsection, we briefly explain how ABH model fits into the previous framework.

Environment: In the ABH model, the economy is populated by a continuum of households, denoted by $I := [0, 1]$. There are two objects: a perishable consumption good and a durable capital shock. There are no stochastic aggregate states.⁵ Each household has utility over

⁵In the notation of Section 2, z_t is a constant, i.e., the drift and volatility in equation (2.1) vanish.

consumption goods of $u(c) = c^{1-\gamma}/(1-\gamma)$, where $\gamma \in (0, 1)$. Each household has two idiosyncratic states: their wealth, a_t^i , and their labour endowment, y_t^i , which can take on one of two values, $y_1 < y_2$. Agents sell their labour endowment in a competitive market for wage, w_t , and they can save or borrow in a competitive market for rate, r_t . In the notation of Section 2, households choose consumption, c_t^i , and their idiosyncratic state vector, $x_t^i = [a_t^i, y_t^i]$, then evolves according to:

$$dx_t^i = d \begin{bmatrix} a_t^i \\ y_t^i \end{bmatrix} = \begin{bmatrix} s(a_t^i, y_t^i, c_t^i, r_t, w_t) \\ 0 \end{bmatrix} dt + \begin{bmatrix} 0 \\ \tilde{y}_t^i - y_t^i \end{bmatrix} dJ_t^i$$

where \tilde{y}_t^i is the complement of y_t^i and the agent's saving function is given by:

$$s(a, y, c, r, w) = wy + ra - c.$$

Each agent is subject to a borrowing constraint that is typically written as the hard constraint that $a_t^i \geq \underline{a}$. However, to make the problem more tractable for the neural network, we follow [Brzoza-Brzezina et al. \(2015\)](#) and introduce a penalty function ψ at the left boundary, replacing the agent flow utility by:

$$\tilde{u}(a_t, c_t) = u(c_t) + \mathbf{1}_{a_t \leq \underline{a}} \psi(a_t)$$

The penalty function we use here is the quadratic: $\psi(a) = -\frac{1}{2}\kappa(a - \underline{a})^2$ where κ is a positive constant. The population density, g_t , is the density across the idiosyncratic states $\{a_t^i, y_t^i\}$ at time t .

There is a production technology, which is controlled by a representative firm. The firm can produce consumption goods according to the production function:

$$Y_t = F(K_t, L_t)$$

where K_t is the capital hired at time t , L_t is the labour hired at time t , and F is the CRS production function $F(K, L) = e^{\bar{Z}} K^\alpha L^{1-\alpha}$. Consumption goods can be converted into capital at 1 – 1 ratio. Capital depreciates at rate δ .

Firm optimization and market clearing implies that, given g_t , the prices r_t and w_t solve:

$$\begin{aligned} r_t &= \partial_K F(K_t, L_t) - \delta, & w_t &= \partial_L F(K_t, L_t), \\ K_t &= \sum_{j \in \{1, 2\}} \int_{\underline{a}}^{\infty} a g_t(a, y_j) da & L &= \sum_{j \in \{1, 2\}} \int_{\underline{a}}^{\infty} y_j g_t(a, y_j) da. \end{aligned} \tag{4.1}$$

Since the prices are implicitly a function of \bar{g} , i.e. of first moments of g , we write them as $r(\bar{g}_t)$ and $w(\bar{g}_t)$. In the notation of Section 2, $Q(\bar{g}) = [r(\bar{g}), w(\bar{g})]$. This is equation that connects the distribution of agents to market prices and so to the household optimizing

behaviour.

Master equation: Let $c^*(a, y, g)$ denote the equilibrium optimal household control. Then, the master equation for the ABH model is given by the following:

$$0 = (\mathcal{L}V)(a, y, g) = (\mathcal{L}^h V)(a, y, g) + (\mathcal{L}^g V)(a, y, g)$$

where the operators \mathcal{L}^h and \mathcal{L}^g are defined by:

$$\begin{aligned} (\mathcal{L}^h V)(a, y, g) &:= -\rho V(a, y, g) + u(c^*(a, y, g)) + \mathbf{1}_{a \leq \underline{a}} \psi(a) \\ &\quad + \partial_a V(a, y, g) s(a, y, c^*(a, y, g), r(\bar{g}), w(\bar{g})) \\ &\quad + \lambda(y)(V(a, \tilde{y}, g) - V(a, y, g)) \\ (\mathcal{L}^g V)(a, y, g) &:= \int_{-\infty}^{\infty} \frac{\partial V}{\partial g}(a, y, g)(b) (\lambda(\tilde{y})g(b, \tilde{y}) - \lambda(y)g(b, y)) db \\ &\quad + \int_{-\infty}^{\infty} \partial_b \frac{\partial V}{\partial g}(a, y, g)(b) s(a, y, c^*(a, y, g), r(\bar{g}), w(\bar{g})) g(b, y) db, \end{aligned}$$

where $r(\bar{g})$ and $w(\bar{g})$ solve the system of equations (4.1), the optimal control satisfies the following FOC:

$$\partial_a V(a, y, g) = u'(c^*(a, y, g)).$$

Since the optimal control is a function of the $\partial_a V(a, y, g)$, it will be more convenient to solve the master equation for the partial derivative, which we denote by $W(a, y, g) = \partial_a V(a, y, g)$. This function is solution to the following PDE:

$$0 = (\mathfrak{L}W)(a, y, g) = (\mathfrak{L}^h W)(a, y, g) + (\mathfrak{L}^g W)(a, y, g)$$

where the operators \mathfrak{L}^h and \mathfrak{L}^g are defined by:

$$\begin{aligned} (\mathfrak{L}^h W)(x, g) &:= (r(\bar{g}) - \rho)W(a, y, g) + \mathbf{1}_{a \leq \underline{a}} \psi'(a) \\ &\quad + \partial_a W(a, y, g) s(a, y, c^*(a, y, g), r(\bar{g}), w(\bar{g})) \\ &\quad + \lambda(y)(W(a, \tilde{y}, g) - W(a, y, g)) \\ (\mathfrak{L}^g W)(x, g) &:= \int_{-\infty}^{\infty} \frac{\partial W}{\partial g}(a, y, g)(b) (\lambda(\tilde{y})g(b, \tilde{y}) - \lambda(y)g(b, y)) db \\ &\quad + \int_{-\infty}^{\infty} \partial_b \frac{\partial W}{\partial g}(a, y, g)(b) s(a, y, c^*(a, y, g), r(\bar{g}), w(\bar{g})) g(b, y) db \end{aligned} \tag{4.2}$$

with the FOC:

$$W(a, y, g) = u'(c^*(a, y, g)).$$

In the next sections, we solve this master equation numerically. The parameters that we use in numerical experiments are in Appendix C.1.

4.2 Details on the Finite Agent Approximation

We start with the finite agent approximation. In this case, we replace the distribution by the positions of the agents:

$$\hat{g} = \{(a^i, y^i)\}_{i \leq I}$$

where $I = 41$ agents. The master equation operators can be written as:

$$\begin{aligned} (\hat{\mathcal{L}}^h W)((a^i, y^i), \hat{g}) &= + (r(\hat{g}) - \rho)W((a^i, y^i), \hat{g}) + \psi_a(a^i) + s_i \partial_a W((a^i, \tilde{y}^i), \hat{g}) \\ &\quad + \lambda(y^i) (W((a^i, \tilde{y}^i), \hat{g}) - W((a^i, y^i), \hat{g})) \\ (\hat{\mathcal{L}}^g W)((a^i, y^i), \hat{g}) &= \sum_{j \neq i} s_j ((a^j, y^j), \hat{g}) \partial_{a^j} W((a^i, y^i), \hat{g}) \\ &\quad + \lambda_j(y^j) (W((a^i, y^i), \{(a^j, \tilde{y}^j), \hat{g}^{-j}\}) - W((a^i, y^i), \hat{g})) \end{aligned}$$

We found that it was customize the generic Algorithm 1 in the following ways to solve the Aiyagari model.

Choice of Neural Network (Step 1): We use a fully connected feed-forward neural network with 5 layers and 64 neurons per layer. We use a tanh activation function between layers and an ELU activation at the output level. We initialize the neural network by pre-training it to have a log shape in a . We tried setting up the network with and without the imposition of symmetry suggested by [Kahou et al. \(2021\)](#) and [Han et al. \(2021\)](#) but ultimately found that the solution was more robust without the symmetry.

Sampling Approach (Step 3a): We sample points of the form $\{(a^i, y^i), \{a^j, y^j\}_{j \in (I-1)}\}$. For the idiosyncratic variable, a^i , we sample using an active sampling technique similar to those developed by [Gopalakrishna \(2021\)](#) and [Lu et al. \(2021a\)](#). We divide the state space for $a \in [0, 20]$ in a collection of regions. For each iteration of the algorithm, we calculate the average loss in each region. We then add additional points to the regions with the largest loss. We use the active sampling approach because it allows us to actively learn where the algorithm is having trouble minimizing the loss function. The interval $[\underline{a}, \bar{a}]$ is evenly partitioned into 2^4 subintervals. We do active sampling after 2,000 of epochs to make it efficient. We calculate the residual error in each subinterval, find where it is the largest and add $2^4, 2^3, 2^2$ points to it, its nearest, and second nearest neighboring subintervals. For sampling the population of agents, $\{a^j, y^j\}_{j \in (I-1)}$, we generate a random interest rate, then generate a random distribution of agents and scale their individual wealth so the equilibrium interest rate is the randomly drawn interest rate. The interest rate r is drawn from a uniform distri-

bution $U[r_{lb}, r_{rb}]$, $r_{lb} = -0.05$, $r_{rb} = 0.05$. This is quite robust to different model parameters.

Loss Function Specification (Step 3b): In order to help the neural network to maintain monotonicity and have non-complex number consumption from the optimality condition $u'(c^*) = W$, we introduced a penalty for having $\partial_a V = W$ less than a lower bound $\underline{W} > 0$.

4.3 Discrete State Space Approximation

We now turn to the second approximation method. Algorithm 1 is implemented in the following way.

Choice of Neural Network (Step 1): We approximate $W = \partial_a V$ by a neural network $\hat{W}(\cdot; \theta_W)$ with parameters θ_W . For more stability of the learning process, we also approximate c^* by another neural network, $c^*(\cdot; \theta_c)$ with parameters θ_c . For both networks, we use an architecture similar to the one proposed by [Sirignano and Spiliopoulos \(2018\)](#). The main differences are two-fold. On the last layer, we use an activation function ensuring that the output is always positive. Furthermore, the first layers of the neural network realize an embedding of the distribution \hat{g} in a space of lower dimension.

Sampling Approach (Step 3a): We sample points of the form (a, y, \hat{g}) where (a, y) follows a uniform distribution over $[\underline{a}, \bar{a}] \times \{y_1, y_2\}$ and \hat{g} is obtained as a randomly perturbed version of the equilibrium steady state \hat{g}_{ss} . More precisely, we take $\hat{g}_{i,j} = \frac{\omega_{i,j} \hat{g}_{ss,i,j}}{\sum_{i=1}^{N_x} \omega_{i,j} \hat{g}_{ss,i,j}}$ for $j = 1, 2$, where the $\omega_{i,j}$ are i.i.d. with uniform distribution over an interval of the form $[1 - d_g, 1 + d_g]$, with $d_g \in (0, 1)$. Thanks to the fact that we deal with the boundary condition through a penalty, we do not need to sample points on the boundary, nor to evaluate the error \mathcal{E}^b .

Loss Function Specification (Step 3b): To compute the error \mathcal{E}^e , the operators $\hat{\mathcal{L}}^h$ and $\hat{\mathcal{L}}^g$ approximating the operators introduced in (4.2) are represented as follows. $\hat{\mathcal{L}}^h \hat{W}(a, y, \hat{g})$ is obtained by evaluating the neural network for \hat{W} and by computing its partial derivative with respect to a using automatic differentiation. $\hat{\mathcal{L}}^g \hat{W}$ is obtained by taking the product of the gradient of \hat{W} with respect to \hat{g} , which is computed using automatic differentiation, and $\hat{\mu}^{\hat{g}}$, which is computed as follows. The ODE (3.1) which describes the evolution of \hat{g} is obtained by using a finite-difference scheme proposed by [Achdou et al. \(2017\)](#). The right-hand side is written in the form $\hat{\mu}_{\hat{g}}(\hat{g}_t) = A^\top \hat{g}_t$, where A^\top is a matrix which approximates spatial derivatives by finite differences. It relies on an upwind scheme for more stability. On top of this error, we also add an extra term that penalizes the neural network when $\partial_a \hat{W}$ is positive. This extra term encourages $\hat{W} = \partial_a \hat{V}$ to be a decreasing function of a , which is consistent with the fact that \hat{V} should be a concave function of a , for any given (y, \hat{g}) .

After each gradient step to update the parameters θ_W of the neural network for \hat{W} , we update the parameters θ_c of the neural network for c^* by minimizing a Monte Carlo

approximation of $|c^*(a, y, \hat{g}; \theta_c) - (u')^{-1}(\hat{W}(a, y, \hat{g}; \theta_W))|^2$.

4.4 Comparison to Finite Difference Solution

In this section, we compare the output from our neural network solution to the output from our a finite difference solution to the Aiyagari model. We first compare the solution at the steady state and then compare the solution along transition paths following MIT shocks.

4.4.1 Steady State Comparison

Figure 1 plots the steady state consumption policy rule, derivative of the value function, probability density function (pdf), and cumulative distribution function (cdf) for the solutions from the finite agent code, the finite state space code, and finite difference code. As can be seen, the neural network solutions align very closely to the finite difference solution. The loss is:

	Master equation loss	MSE(NN, FD)
Finite Agent NN	3.135×10^{-5}	4.758×10^{-5}
Discrete State Space NN	2.548×10^{-4}	1.590×10^{-2}

Table 1: Neural Nets’ results for solving Master Equations. The Master equation loss is the mean squared error of residuals. MSE(NN,FD) is the mean squared difference of consumption solved by neural network and finite difference. The training loss vs iterations are available in Appendix D.

4.4.2 Comparison to Finite Difference Solution: Transition Dynamics

We can also compare the transition path. An important advantage of having a global solution is that we can solve for the transition path without a “shooting algorithm”, as is commonly done in the finite difference literature. This is advantage because shooting algorithms are often unstable, particular for systems with a large number of prices that require complicated guess for the price path. Instead, with a full solution to the master equation, we can solve the Kolmogorov Forward Equation (KFE) directly as time-dependent consumption is a function of density $g(a, y)$.

$$\frac{\partial g(a, y)}{\partial t} = -\frac{\partial}{\partial a}(s(a, y, g)g(a, y)) + \lambda(g(a, \tilde{y}) - g(a, y)) \equiv \mathcal{A}^T g$$

where \mathcal{A} is transition matrix. We illustrate this for the finite agent approximation. The iterative procedure for solving the KFE in this case is summarized in Algorithm 2 below. We expand on the different approaches for calculating the transition path in Appendix B.3.

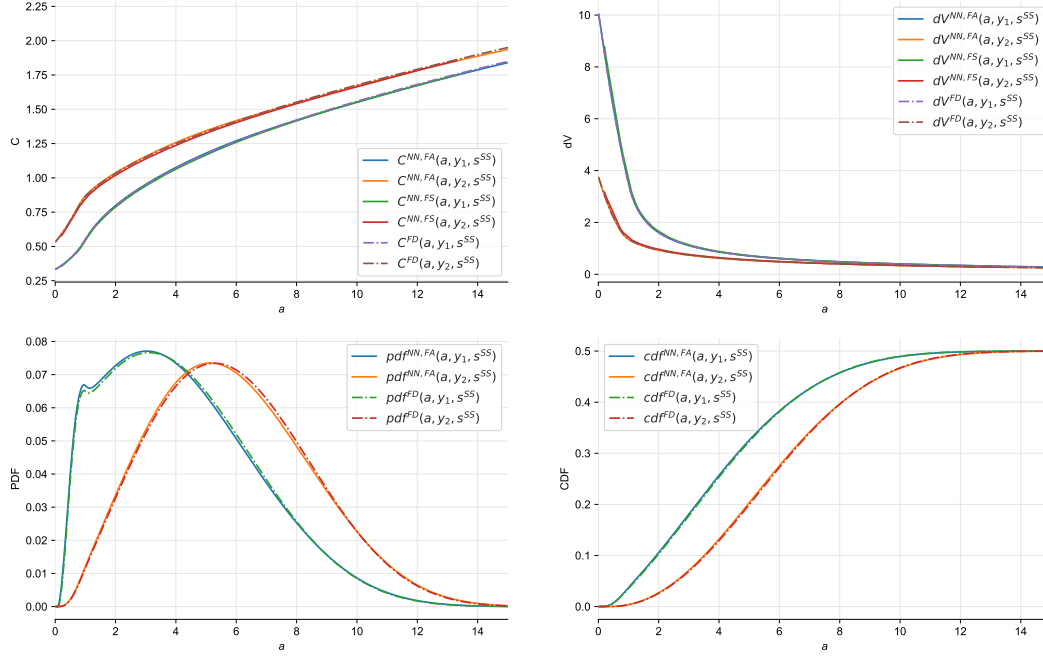


Figure 1: Comparison between neural network and finite difference solutions for the Aiya-gari model. The top left plot shows the consumption policy rule. The top right shows the derivative of the value function, the bottom left shows the pdf, and the bottom right shows the cdf. The superscript NN, FA refers to the finite agent neural network code, the superscript NN, FS refers to the finite state space neural network code, and the superscript FD refers to the finite difference code.

We compare the neural network and finite difference transition paths in Figure 2 below. We discuss how the finite difference solution is computed in Appendix B.3. In this numerical experiment, we train our neural network at $Z = 0$ and we start from an economy in its steady state with productivity $Z_t = -0.10$ for $t = 0$. At $t = 0^+$, an unexpected positive productivity shock brings Z from -0.10 to $Z_t = 0$ permanently. We solve distributional dynamics by Algorithm 2, and use steady state at $Z = -0.10$ as the initial condition. We plot the percentage change of capital, capital return and wage evolution first row and second row of Figure 2. The difference between neural net's transition paths and finite difference's transition paths are less than 0.1%. The lower panels of Figure 2 compares neural network and finite difference probability density at time $t = 15$ and $t = 30$.

5 Example: Uninsurable Income Risk and TFP Shocks

A canonical extension of the previous model is the [Krusell and Smith \(1998\)](#) model, which introduces TFP shocks. This model is often used as a test model for new numerical techniques.

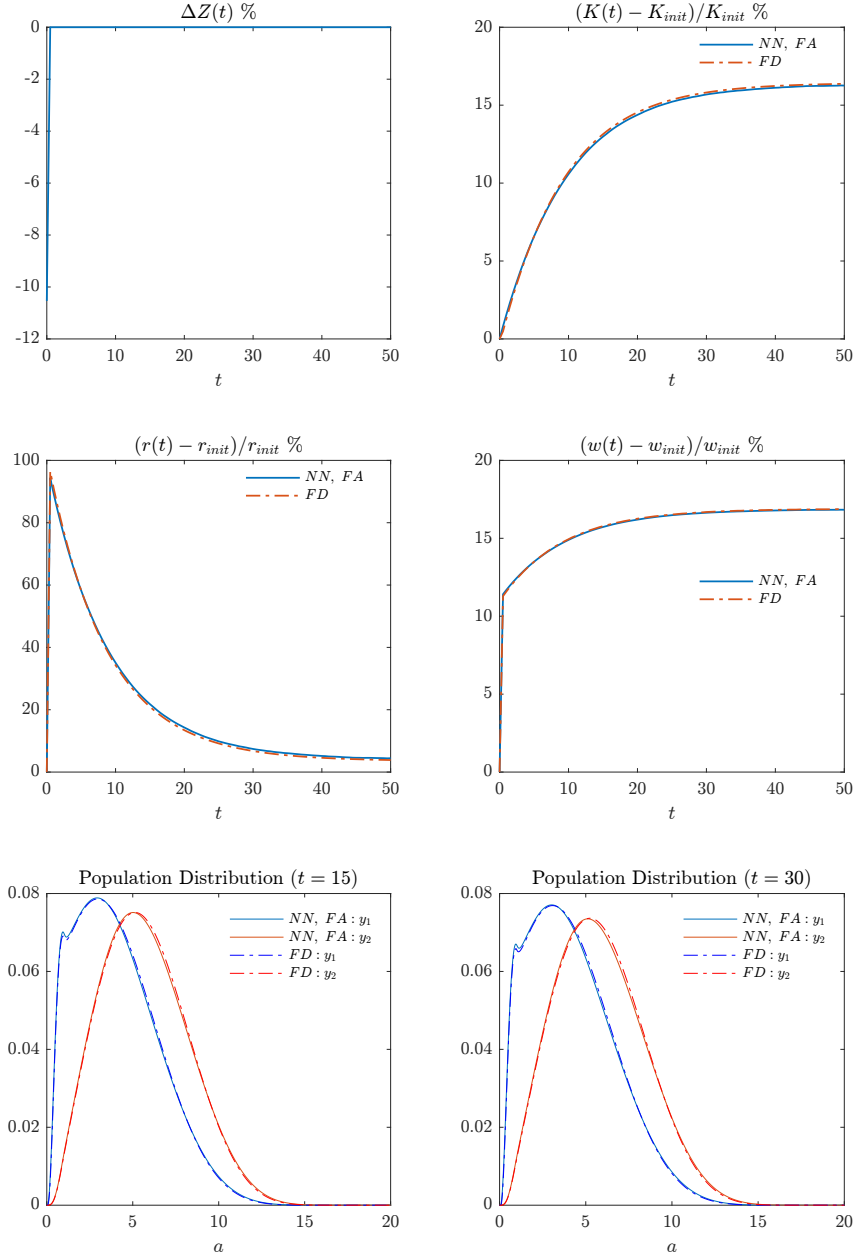


Figure 2: Comparison between neural network and finite impulse response for the Aiyagari model. The top left plot is the TFP shock path, the top right panel is the aggregate relative capital change, the middle left panel plots the relative capital return change, and the middle right panel plots the relative wage change. The bottom left and bottom right are snapshots of probability density at $t = 15$ and $t = 30$. *NN, FA* refers to the finite agent neural network code, and the *FD* refers to the finite difference code. Subscript *init* is the initial state at the steady state $Z = Z(0)$.

Algorithm 2: Finding Transition Path by Neural Network: Aiyagari case

1. Sample N agents from distribution $g_t(a, y)$ at time t .
 2. Given other agents' state s_{-i} , calculate the consumption $c(a, y, s_{-i})$, equilibrium capital return $r(s_{-i})$ and wage $w(s_{-i})$. Then construct \mathcal{A} by finite difference.
 3. Repeat 1. and 2. for N_{sim} times, then take the average: $\bar{\mathcal{A}} = \frac{1}{N_{sim}} \sum_{i=1}^{N_{sim}} \mathcal{A}_i$
 4. Update $g_t(a, y)$ by implicit method: $g_{t+dt} = (I - \bar{\mathcal{A}}^T dt)^{-1} g_t$
-

Environment Changes: The model is the same as the previous section, except that now the firm can produce consumption goods according to the production function:

$$Y_t = e^{Z_t} F(K_t, L_t)$$

where Z_t is the aggregate productivity, which follows an Ornstein-Uhlenbeck process:

$$dZ_t = \eta(\bar{Z} - Z_t)dt + \sigma dB_t^0$$

For this model, g_t now denotes the population density across $\{a_t^i, y_t^i\}$ at time t , given a filtration, \mathcal{F}_t^0 , generated by the sequence of aggregate shocks. Given g_t and Z_t , the prices r_t and w_t solve (and so are implicitly function of \bar{g}_t and Z_t):

$$\begin{aligned} r_t &= e^{Z_t} \partial_K F(K_t, L) - \delta, & w_t &= e^{Z_t} \partial_L F(K_t, L), \\ K_t &= \sum_{j \in \{1, 2\}} \int_{\underline{a}}^{\infty} a g_t(a, y_j) da & L &= \sum_{j \in \{1, 2\}} \int_{\underline{a}}^{\infty} y_j g_t(a, y_j) da \end{aligned} \quad (5.1)$$

The master equation operators become:

$$\begin{aligned} (\mathcal{L}^h V)(a, y, Z, g) &:= -\rho V(a, y, Z, g) + u(c^*(a, y, Z, g)) + \mathbf{1}_{a \leq \underline{a}} \psi(a) \\ &\quad + \partial_a V(a, y, Z, g) s(a, y, c^*(a, y, Z, g), r(\bar{g}), w(\bar{g})) \\ &\quad + \lambda(y)(V(a, \tilde{y}, Z, g) - V(a, y, Z, g)) \\ &\quad + \partial_Z V(a, y, Z, g) \eta(\bar{Z} - Z) + \frac{1}{2} \sigma^2 \partial_{ZZ} V(a, y, Z, g) \\ (\mathcal{L}^g V)(a, y, Z, g) &:= \int_{\underline{a}}^{\infty} \frac{\partial V}{\partial g}(a, y, Z, g)(b) (\lambda(\tilde{y})g(b, \tilde{y}) - \lambda(y)g(b, y)) db \\ &\quad + \int_{\underline{a}}^{\infty} \partial_b \frac{\partial V}{\partial g}(a, y, g)(b) s(a, y, c^*(a, y, Z, g), r(\bar{g}, Z), w(\bar{g}, Z)) g(b, y) db \end{aligned}$$

where $r(\bar{g}, Z)$ and $w(\bar{g}, Z)$ solve the system of equations (5.1), the optimal control satisfies:

$$\partial_a V(a, y, Z, g) = u'(c^*(a, y, Z, g))$$

5.1 Finite Agent Approximation

We customize the generic Algorithm 1 in the following ways to solve the Krusell-Smith model. We solve a model with 41 agents.

Choice of Neural Network (Step 1): We use a fully connected feed-forward neural network with 5 layers and 64 neurons per layer. We use a tanh activation function between layers. We initialize the neural network by pre-training it to have a exp shape in a . The same as in the Aiyagari model, we find the solution was more robust without symmetry.

Sampling Approach (Step 3a): We sample points of the form $\{(a^i, y^i), \{a^j, y^j\}_{j \in (I-1)}, Z\}$. For idiosyncratic variable, (a^i, y^i) , and population of agents $\{a^k, y^j\}_{j \in (I-1)}$, the sampling method is the same as in section 4.2. We sample aggregate variable Z uniformly from interval $[Z_{min}, Z_{max}]$.⁶ Again, this is quite robust to different model parameters.

Loss Function Specification (Step 3b): In order to help the neural network to maintain monotonicity and have non-complex number consumption from the optimality condition $u'(c^*) = W$, we introduced a penalty for having $\partial_a V = W$ less than a lower bound $\underline{W} > 0$. To capture curvature in Z dimension, we introduced a penalty for having $d\bar{W}/dZ > 0$, which is equivalent to penalize $dC/dZ < 0$, where C is aggregate consumption.

5.2 Numerical Solution

We solve the Krusell-Smith model using the finite agent method. The error in the master equation is shown in Table 2 below.

Master equation loss	
Finite Agent NN	3.037×10^{-5}

Table 2: Neural Nets' results for solving Master Equations with aggregate shocks.

Unlike for the Aiyagari model, we don't have a clear benchmark for Krusell-Smith model because there is no existing technique that provides an accurate solution to the model with

⁶Theoretically, it is possible that Z_t becomes lower than Z_{min} or greater than Z_{max} . In practice, letting $|Z_{min}/max| = 4\sigma$ is good enough to approximate the economic relevant region. We cannot replace the step of having $r \in [r_{lb}, r_{rb}]$ by having $Z \in [Z_{min}, Z_{max}]$ to let the distribution move around because Z 's scaling effect is weaker. In fact Z 's impact on interest rate is $\Delta r \approx (r + \delta_K)\Delta Z$, which can hardly let the population distribution move around. See more discussions in Section 6.

aggregate shocks. However, we can compare to widely used approximation techniques in the literature. In particular, we compare the approach suggested by [Fernández-Villaverde et al. \(2018\)](#), which uses a neural network to approximate a statistical law of motion rather than developing the fully global solution.

We compare to [Fernández-Villaverde et al. \(2018\)](#) by computing sample paths from both solution approaches. We describe how we generate sample paths for the neural network solution in Algorithm 3 below. Essentially, we draw a series of productivity shocks from the Ornstein-Uhlenbeck process: $dZ_t = \eta(\bar{Z} - Z_t)dt + \sigma dW_t$ and then evolve the population distribution by adding Z_t to Algorithm 2 for the ABH model described in section 4.4.2.

Algorithm 3: Finding Transition Path by Neural Network: Krusell Smith case

1. Sample dB_t from normal distribution $N(0, dt)$, construct TFP shock path by:
 $Z_t = Z_{t-dt} + \eta(\bar{Z} - Z_{t-dt}) + \sigma dB_t$.
 2. Sample N agents from distribution $g_t(a, y)$ at time t .
 3. Given other agents' state s_{-i} , calculate the consumption $c(a, y, s_{-i}, Z_t)$, equilibrium capital return $r(s_{-i}, Z_t)$ and wage $w(s_{-i}, Z_t)$. Then construct \mathcal{A} by finite difference.
 4. Repeat 2. and 3. for N_{sim} times, then take the average: $\bar{\mathcal{A}} = \frac{1}{N_{sim}} \sum_{i=1}^{N_{sim}} \mathcal{A}_i$
 5. Update $g_t(a, y)$ by implicit method: $g_{t+dt} = (I - \bar{\mathcal{A}}^T dt)^{-1} g_t$
-

Figure 3 shows the comparison between our neural network solution and [Fernández-Villaverde et al. \(2018\)](#) for a particular path of productivity shocks. The upper-left panel shows the draw from the Ornstein-Uhlenbeck process: $dZ_t = \eta(\bar{Z} - Z_t)dt + \sigma dB_t$. The upper left compares the evolution of capital stock. The middle plots compares the evolution of prices. The bottom plots compare the evolution of the population. As can be seen in figure 3, we get a similar path for aggregate capital stock, interest rates, wage rates, and the population distribution.

In figure 4, we generate multiple random paths for TFP, Z_t , and show the evolution of our Neural Network solution and solution in [Fernández-Villaverde et al. \(2018\)](#) in a “fan chart” that displays percentiles for the evolution of the population. In particular, we generate 1,000 TFP paths starting from $Z_0 = 0$ and calculate the corresponding aggregate capital evolution paths. We collect capital at different time t , sort to get the quantile $K_t^Q = Q(p)$ and plot the time series as in a fan chart.

6 Practical Lessons

Although the deep learning algorithm is straightforward to describe, we find that implementing it successfully can be tricky. In this section, we collect some lessons that we believe

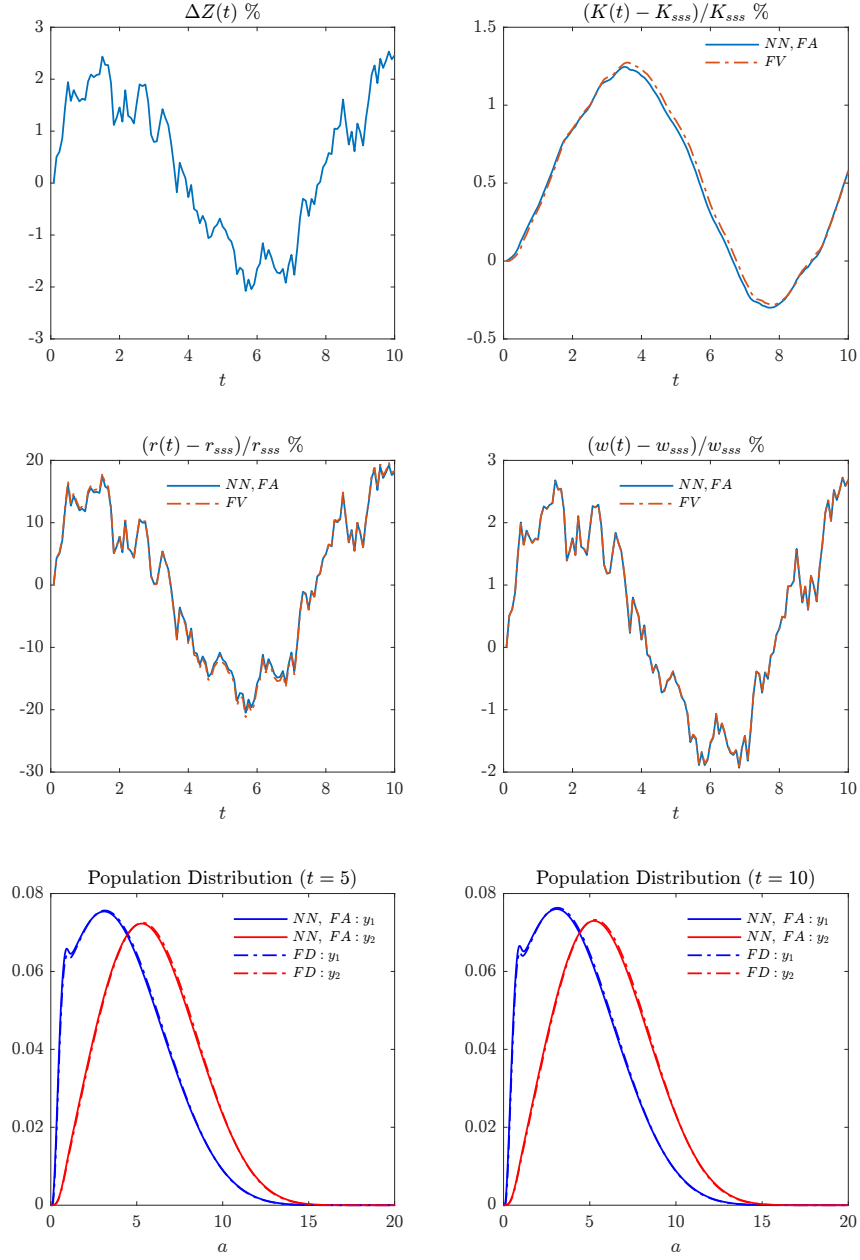


Figure 3: Impulse response functions for Krusell-Smith Model. The top left plot is the TFP shock path, the top right panel is the aggregate relative capital change, the middle left panel plots the relative average consumption change, and the middle right panel plots the relative capital return change. The bottom left is relative wage change, and the bottom right is the relative wealth change at different quantiles. *NN, FA* refers to the finite agent neural network and *FV* refers to the result generated from [Fernández-Villaverde et al. \(2018\)](#). Subscript *sss* refers to the stochastic steady state at $Z = 0$.

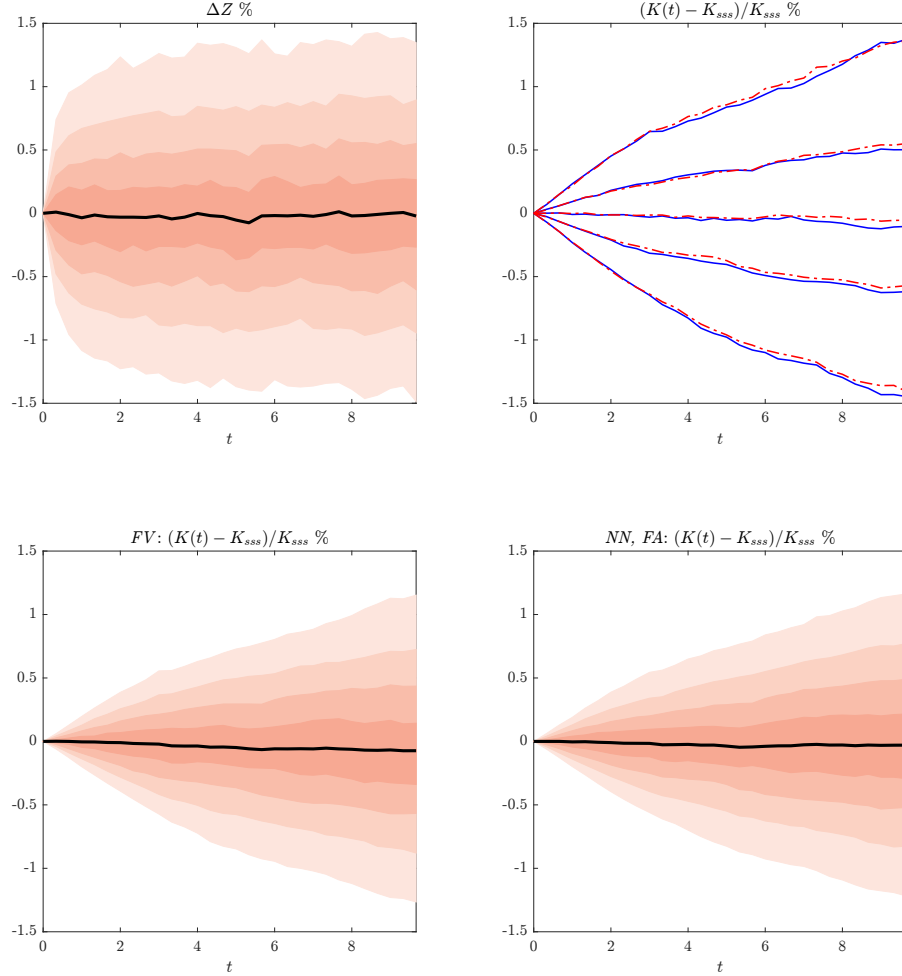


Figure 4: Forecasted aggregate capital dynamics starting from the stochastic steady state (ss) for the Krusell-Smith Model. The top left plot is the fan chart for the TFP shock path, generated from OU process with initial condition $Z_0 = 0$. The bottom left panel and right panel are fan charts (capital quantile) of corresponding responses. The top right panel is the time series plot for relative change in aggregate capital at quantile 10%, 30%, 50%, 70%, 90% (from the lowest to the highest), in which the blue solid lines are generated by neural network solution and the red dashed lines are generated by Fernández-Villaverde et al. (2018). *NN*, *FA* refers to the finite agent technique, and *FV* refers to Fernández-Villaverde et al. (2018)'s technique.

are helpful for using deep learning to solve macroeconomic models.

Lesson 1: Working out the correct sampling approach is very important. A key feature of continuous time methods is that we must specify where in the state space to sample. This is very different to discrete time approaches which typically simulate the economy and so implicitly sample from the ergodic distribution. Like [Gopalakrishna \(2021\)](#), we find that choosing where to sample is both an advantage (because we can focus sampling in interesting, rarely visited regions of the state space with complicated curvature) but also a difficulty (because it can be hard to know where to sample). Many of our initial problems were resolved by adjusting how we were sampling the finite dimensional approximation to the distribution. In particular, we found it is essential to have significant variation in distribution so that the neural network can learn where there is curvature in the problem.

Lesson 2: Neural networks have difficulty dealing with inequality constraints. The [Achdou et al. \(2022\)](#) formulation of the [Aiyagari \(1994\)](#) model is written with a hard lower boundary that $a_t \geq \underline{a}$, which leads to an inequality boundary condition at \underline{a} and a mass point in the distribution at \underline{a} . This is convenient for the finite difference solution technique because the inequality constraint does not impact the tractability of the upwind scheme. However, it causes problems for the neural network approach because the inequality constraint is too “easy” to satisfy, in the sense that many levels of the value function satisfy the constraint. Placing a low weight on the lower boundary error leads to solutions with the correct curvature but the wrong level. Placing a high weight on the lower boundary error leads to solutions with the right level but inaccurate curvature. So, there seems to only be a very small subset of weights that lead to accurate results for inequality boundary conditions. Ultimately, we find that replacing the hard constraint by a soft constraint and utility penalty makes the solution much more robust.

Lesson 3: Enforcing shape constraints is very important. We find that the neural network is likely to converge to “cheat solutions” that approximately solve the differential equation by setting derivatives to zero. There are number of ways to help with this: (i) imposing loss functions on the curvature of the value function to enforce the correct shape (e.g. penalizing non-monotonicity or non-concavity), (ii) pre-training the neural network to match an initial guess known properties of the value function, and (iii) sampling from a sufficiently large part of the state space that the neural network realizes there is curvature in all dimensions.

Lesson 4: Mean squared errors can be misleading. We found that even if mean squared training errors are in the order of 10^{-2} or 10^{-3} , the neural network can give policy rules that are highly inaccurate. This suggests that overfitting can be a large problem for neural network solutions to PDEs and so choosing the right “cross validation” sample is important. It also suggests that the threshold for convergence for neural network solutions to contin-

uous time models might need to be higher than for other techniques, such as finite difference.

Lesson 5: Start with a simple model to tune hyperparameters. A benefit and cost with using neural networks is that they are very flexible approximations. This means that a difficult and time consuming part of training a neural network model is finding appropriate hyperparameters. We find that it is helpful to start with a simple model that can be solved with finite difference (e.g. a version of the model without aggregate shocks) and then tune the hyperparameters to make sure that the neural network approximation closely matches the finite difference solution. We can then introduce aggregate shocks starting with a set of hyperparameters that are good at capturing many features of the distribution. In the language of machine learning, we suggest deploying a combination of supervised learning on a simple version of the problem and then unsupervised learning on the more complicated problem.

7 Conclusion

This paper proposed a new global solution algorithm for continuous time heterogeneous agent economies with aggregate shocks. We demonstrated our algorithm by solving two canonical models in the macroeconomics literature: the [Aiyagari \(1994\)](#) model and the [Krusell and Smith \(1998\)](#) model. However, this is only the beginning of what is possible with this technique. Future work can deploy this solution technique to solve high dimensional economic models with non-linearities and aggregate shocks.

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A Additional Theoretical Results for Section 2

In this section of the appendix, we outline additional theoretical results relating to section 2 of the main text. We first outline a more general version of the economic environment for which our solution techniques are appropriate.

A.1 General Economic Model

In this section of the appendix, we outline the more general version of the economic model for which the techniques in this paper are appropriate. Relative to Section 2, we generalize the model among three dimensions. First, we allow the aggregate state z_t^i to be of arbitrary dimension and driven by both Brownian and Poisson noise. Second, we allow the idiosyncratic state x_t^i to be driven by more than one Poisson processes. Third, we allow the control c_t^i to be multi-dimensional and to affect also the diffusion and jump terms of x_t^i . These generalizations complicate the notational burden required to present the model but do not pose any additional conceptual challenges for our solution algorithms. We present here only the aspects of the model that change relative to Section 2.

Aggregate State Dynamics: There is an exogenous aggregate state vector, $z_t \in \mathcal{Z}$, of arbitrary (finite) dimension. z_t evolves according to:

$$dz_t = \mu^z(z_t)dt + \sigma^z(z_t)dB_t^0 + \gamma^z(z_t)dJ_t^0, \quad (\text{A.1})$$

where B^0 denotes a common N^{B^0} -dimensional Brownian motion process and J^0 denotes a common N^{J^0} -dimensional Poisson jump process. The aggregate history \mathcal{F}_t^0 is defined as the filtration generated by B^0 and J^0 . We let $\nu^n(z)$, $n \leq N^{J^0}$, denote the rate at which the n -th component of the Poisson jump shock arrives given aggregate state z .

Idiosyncratic State Dynamics: Each agent, $i \in I$, has an idiosyncratic state vector, $x^i \in \mathcal{X}$, that evolves according to:

$$dx_t^i = \mu^x(c_t^i, x_t^i, z_t, q_t)dt + \sigma^x(c_t^i, x_t^i, z_t, q_t)dB_t^i + \gamma^x(c_t^i, x_t^i, z_t, q_t)dJ_t^i, \quad (\text{A.2})$$

where $c_t^i \in \mathcal{C}$ is a control vector chosen by the agent, $q_t \in \mathcal{Q}$ is a collection of aggregate prices in the economy, B_t^i denotes an idiosyncratic N^{B^i} -dimensional Brownian motion process, and J^i denotes an idiosyncratic N^{J^i} -dimensional Poisson jump process. We let $\lambda^n(x, z)$, $n \leq N^{J^i}$, denote the rate at which the n -th component of the Poisson jump shock arrives given idiosyncratic state x and aggregate state z .

Agent Problem: Let $u(c_t^i)$ denote the flow utility from choosing the state vector c_t^i . The agent problem is in complete analogy to Section 2. Given their belief about prices q , agent

i chooses their control process, c^i , to solve:

$$\begin{aligned} V(x_0^i, z_0) &= \max_{c^i} \mathbb{E}_0 \left[\int_0^\infty e^{-\rho t} u(c_t^i) dt \right] \\ \text{s.t. } & \text{(A.1), (A.2), } \Phi(x_t^i) \geq 0. \end{aligned} \quad (\text{A.3})$$

Equilibrium: The definition of equilibrium is word by word identical to the definition given in Section 2 in the main text.

KFE: Because dB_t^0 and dJ_t^0 do not directly enter the idiosyncratic state evolution (A.2), the KFE is precisely as in the main text, equation (2.4).

Master Equation: The master equation can again be written in the form

$$(\mathcal{L}V)(x, z, g) = 0$$

with a differential operator $(\mathcal{L}V)(x, z, g) := (\mathcal{L}^h V + \mathcal{L}^g V)(x, z, g)$.

The distribution portion $\mathcal{L}^g V$ of this operator takes precisely the same form as in the main text. It is for this reason that the numerical solution approach readily generalizes to the model version presented here.

The optimization problem portion $\mathcal{L}^h V$ of the differential operator is structurally similar to the main text but requires slightly heavier notation:

$$\begin{aligned} (\mathcal{L}^h V)(x, z, g) &:= -\rho V(x, z, g) + u(c^*(x, z, g)) \\ &+ D_x V(x, z, g) \mu^x(c^*(x, z, g), x, z, Q(z, \bar{g})) + D_z V(x, z, g) \mu^z(z) \\ &+ \frac{1}{2} \text{tr} \{ \Sigma^x(c^*(x, z, g), x, z, Q(z, \bar{g})) D_x^2 V(x, z, g) \} + \frac{1}{2} \text{tr} \{ \Sigma^z(z) D_z^2 V(x, z, g) \} \\ &+ \sum_{n \leq N^{J_i}} \lambda^n(x, z) (V(x + \gamma^{xn}(c^*(x, z, g), x, z, Q(z, \bar{g})), z, g) - V(x, z, g)) \\ &+ \sum_{n \leq N^{J_0}} \nu^n(z) (V(x, z + \gamma^{zn}(z), g) - V(x, z, g)) \end{aligned}$$

Here, the notation is in analogy to the main text: $\Sigma^x(\cdot) = \sigma^x(\cdot) (\sigma^x(\cdot))^T$, $\Sigma^z(\cdot) = \sigma^z(\cdot) (\sigma^z(\cdot))^T$, and $c^*(x, z, g)$ denotes the (recursive) optimal control that solves the agent problem (A.3).

B Aiyagari Model

B.1 Parameters for Aiyagari Model

Parameter	Symbol	Value
Capital share	α	1/3
Depreciation	δ	0.1
Risk aversion	γ	2.1
Discount rate	ρ	0.05
Mean TFP	\bar{Z}	0.00
Reversion rate	η	0.50
Volatility of TFP	σ	0.01
Transition rate (1 to 2)	λ_1	0.4
Transition rate (2 to 1)	λ_2	0.4
Low labor productivity	y_1	0.3
High labor productivity	y_2	$1 + \lambda_2/\lambda_1(1 - y_1)$
Borrowing constraint	\underline{a}	10^{-6}
Maximum of asset	\bar{a}	20.0
Penalty Function	$\psi(a)$	$-\frac{1}{2}\kappa(a - a_{lb})^2$
Penalty parameters	a_{lb}	1.0
Penalty parameters	κ	3.0

B.2 Penalty Function Approximation

In this section, we compare the finite difference solutions with the hard and soft boundary constraints. The comparisons are summarized in Figure 5.

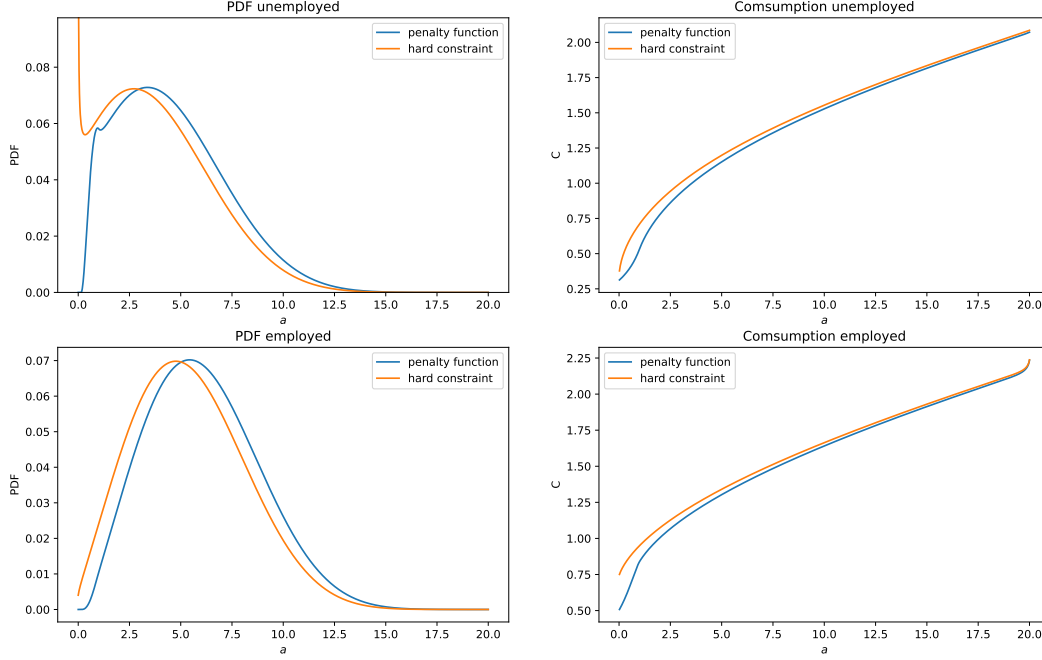


Figure 5: Comparison between penalty function approach and hard constraint in finite different exercise. $\psi(a) = -\frac{1}{2}\kappa(a - a_{lb})^2$, $a_{lb} = 1.0$, $\kappa = 3.0$.

B.3 Comparison of Techniques for Calculating Transition Dynamics without Aggregate Shock

After we obtained the neural network approximation of the value function in the master equation, $V(a_i, y_i, s_{-i})$, we are able to consider the transition path in two ways. The first way is to consider an evolving finite agent economy, the initial asset and employment status are sampled from the initial distribution: $g_0(a, y)$. This method gives the transition path of return on capital/labor directly. To solve the Kolmogorov Forward Equation, the key is to figure out $\mu(a, y) = ra + wy - c$, which can be approximated by the consumption policy. The algorithm can be summarized as follows in a high level.

Algorithm 4: Finding Transition Path by Neural Network

1. Sample N agents from the initial condition $g_0(a, y)$.
 2. Calculate the consumption, then find the transition path for every asset position.
 3. After we've obtained c_t, a_t, r_t , solve $g_t(a, y)$ by finite difference of the forward equation
-

The second way is to update the distribution by Kolmogorov Forward Equation and

resample agents from the updated distribution accordingly. Recall the forward equation in this economy:

$$\frac{\partial g_t(a, y)}{\partial t} = -\frac{\partial(\mu(a, y)g(a, y))}{\partial a} - \lambda_y g(a, y) + \lambda_{\tilde{y}} g(a, \tilde{y}).$$

In the this finite agents economy, the dynamic of equilibrium return on capital does not require another “guess-verify” loop, as in [Achdou et al. \(2022\)](#). The following algorithm modified to solve the dynamics of Aiyagari Economy.

Algorithm 5: Finding Transition Path by Finite Difference

1. Guess the path of equilibrium interest rate $r^o(t)$, then solve HJB, with terminal condition: $v(a, y, T) = \bar{v}(a, y)$
 2. Solve the policy function $c_t(a, y)$.
 3. Solve the forward equation, with initial condition $g_0(a, y)$.
 4. Calculate the capital held by the whole economy: $\int_a^\infty \sum_{y \in \{0,1\}} ag_t(a, y) = K_t$, then calculate the implied interest rate by $r^n(t) = \partial_K F(\bar{K}_t, L) - \delta$, for every $t \in [0, T]$.
 5. Update the path to be $\lambda r^o(t) + (1 - \lambda)r^n(t)$, repeat 1-4 until $\|r^o - r^n\|_\infty < \epsilon$.
-

Lessons. When using iterative method to find steady state/distributional dynamics in the forward equation, we have:

$$g = (I - \mathcal{A}(g)dt)^{-1}g,$$

which means we essentially have a high dimensional *fixed-point* problem. As we are using simulation to find \mathcal{A} , we know \mathcal{A} may be very kinky because of the sampling error, and the solution is to have N_{sim} very large to smooth $(I - \bar{\mathcal{A}}dt)^{-1}$. Otherwise, the distribution g can be trapped before it reaches the steady state, as shown in Figure 6. Initial distribution with total capital $K_{init} < K_{ss,new}$ will end up with a oscillating, but a steady state with slightly lower aggregate capital. Similarly, initial distribution with total capital $K_{init} > K_{ss,new}$ will end up with a oscillating, but a steady state with a slightly higher aggregate capital.

B.4 Compare neural nets with symmetry and without symmetry

- **Speed:** neural nets with symmetry is 10 times slower than without symmetry.
- **Convergence:** neural nets with symmetry has converged (without batch sampling), but it is extremely hard to converge without symmetry
- **Shape:** nn with symmetry has problem capturing the shape, nn without symmetry has problem capturing convexity for KS calibration.

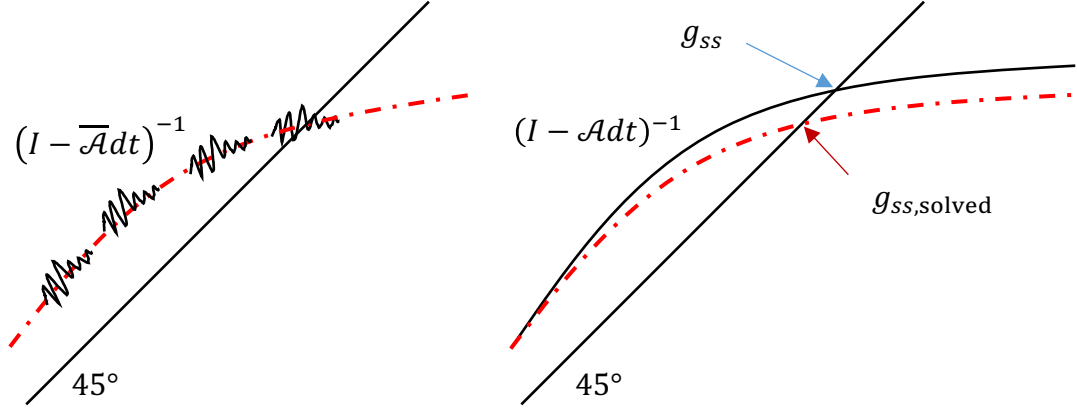


Figure 6: Fixed point problem in finding steady state and distributional dynamics. The fixed point is solved by the cross between $(I - \bar{\mathcal{A}}dt)^{-1}$ and 45° line. The kinky black line on the left panel is the simulated operator $(I - \bar{\mathcal{A}}dt)^{-1}$, and the red dashed line is the smoothed approximation of $(I - \bar{\mathcal{A}}dt)^{-1}$. The concave black solid line on the right panel is the theoretical $(I - \bar{\mathcal{A}}dt)^{-1}$. g_{ss} and $g_{ss,solved}$ are the crossing points between the black solid line and 45° line, the red dashed line and 45° line, respectively.

B.5 Active Learning

To efficiently train our neural net, we use the active learning method to track where losses are larger. We start from a uniform sampler in domain $[\underline{a}, \bar{a}]$. To capture the curvature where penalty function is applied, we add additional points, which is also a hyper parameter can be tuned, to the interval $[\underline{a}, a_{lb}]$. It is crucial to have a better approximation of policies in this subdomain to avoid wrong solutions. In addition, we dynamically adjust the sampler's distribution according to reported losses, similar to [Gopalakrishna \(2021\)](#). We increase the probability density for the subdomains where the loss are larger, and vice versa.

In practice, we partition the domain into N subdomains as in section 2.8 of [Lu et al. \(2021a\)](#). After approximately 2,000 epochs of training, we start to implement active learning. We calculate the master equation's relative residual in each subdomain, and re-weight sampler's distribution accordingly. We find it's better to also increase density to the subdomains close to where the relative residual is the largest. The loss has a steeper drop after we start active learning in the 2000th epoch, as shown in Figure 8.

C Krusell-Smith Model

C.1 Parameters for Krusell-Smith Model

Parameter	Symbol	Value
Capital share	α	1/3
Depreciation	δ	0.1
Risk aversion	γ	2.1
Discount rate	ρ	0.05
Mean TFP	\bar{Z}	0.00
Reversion rate	η	0.50
Volatility of TFP	σ	0.01
Transition rate (1 to 2)	λ_1	0.4
Transition rate (2 to 1)	λ_2	0.4
Low labor productivity	y_1	0.3
High labor productivity	y_2	$1 + \lambda_2/\lambda_1(1 - y_1)$
Borrowing constraint	\underline{a}	10^{-6}
Maximum of asset	\bar{a}	20.0
Penalty Function	$\psi(a)$	$-\frac{1}{2}\kappa(a - a_{lb})^2$
Penalty parameters	a_{lb}	1.0
Penalty parameters	κ	3.0
Drift in O-U Process	η	0.5
Volatility in O-U Process	σ	0.01
Maximum TFP	Z_{max}	0.04
Minimum TFP	Z_{min}	-0.04

D Additional Plots

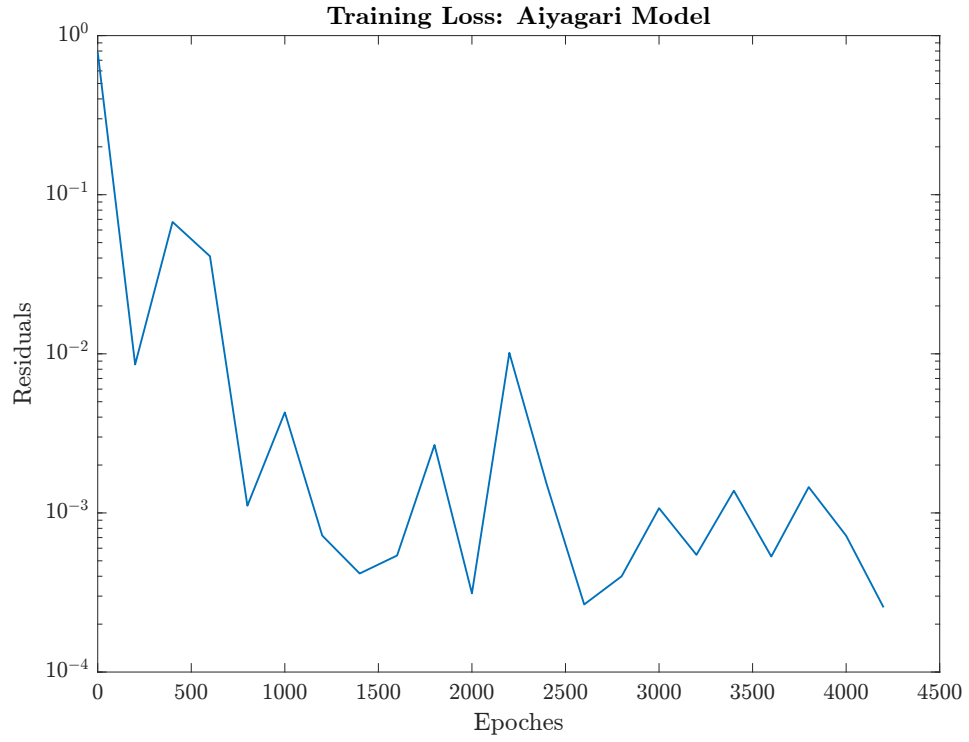


Figure 7: Training Loss vs Iteration Plots (Discrete State Space Method)

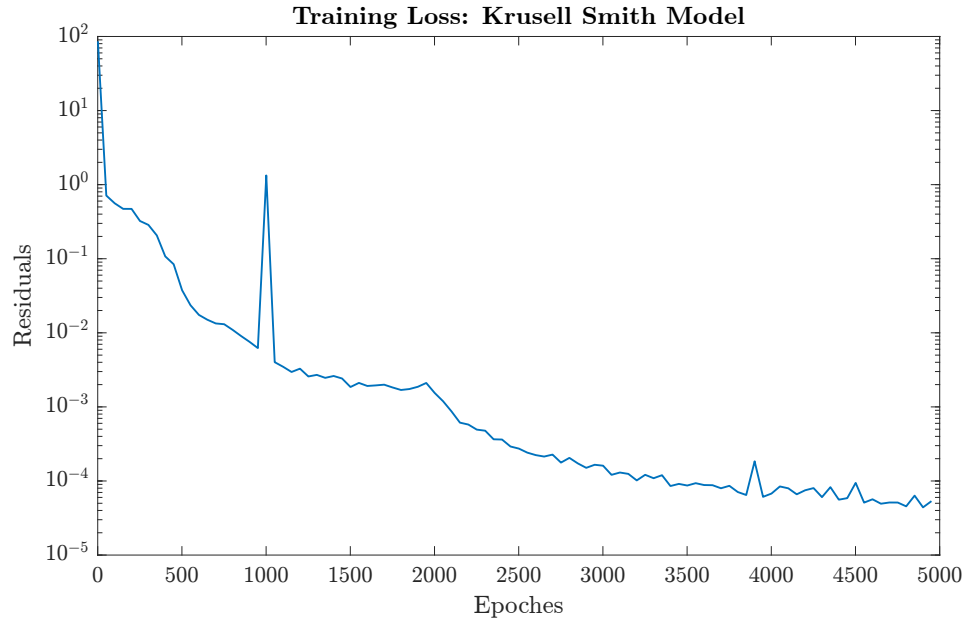


Figure 8: Training Loss vs Iteration Plots (Finite Agent Method)