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DEEP EQUILIBRIUM NETS*

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We introduce deep equilibrium nets (DEQNs)—a deep learning-based method to compute approximate functional rational expectations equilibria of economic models featuring a significant amount of heterogeneity, uncertainty, and occasionally binding constraints. DEQNs are neural networks trained in an unsupervised fashion to satisfy all equilibrium conditions along simulated paths of the economy. Since DEQNs approximate the equilibrium functions directly, simulating the economy is computationally cheap, and training data can be generated at virtually zero cost. We demonstrate that DEQNs can accurately solve economically relevant models by applying them to two challenging life-cycle models and a Bewley-style model with aggregate risk.

1. INTRODUCTION

Heterogeneity between types of agents, such as hand-to-mouth and non hand-to-mouth consumers (see, e.g., Bilbiie, 2008; Debortoli and Galí, 2017; Kaplan et al., 2018), financial frictions (see, e.g., Fernández-Villaverde et al., 2016; Dou et al., 2017, and references therein), such as borrowing constraints, and distributional channels (see, e.g., Krueger et al., 2016) are widely recognized as key ingredients for modern macroeconomic models. However, solving economic models with these features in discrete time remains a challenging and sometimes prohibitive task. Substantial computational challenges arise because of (a) stochasticity, (b)

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very high-dimensional state spaces, (c) strong nonlinearities or kinks in the equilibrium functions, and (d) irregular geometries of the ergodic set of states. In the presence of all of these features, the curse of dimensionality (Bellman, 1961), which is caused by the high-dimensional state space, imposes a substantial roadblock. Computational methods to ameliorate the curse of dimensionality are available for special cases but mostly rely on the absence of at least some of the other features listed above. A model with financial frictions, for instance, often features occasionally binding constraints, which induce kinks in the equilibrium functions. Even methods that are designed to approximate such functions, such sparse grids,¹ fail if the dimensionality of the state space exceeds ~ 20 . Although there are methods that can handle a subset of (a), (b), (c), and (d), there is a need for tractable computational methods that jointly address all four features.²

In this article, we present a novel algorithm based on unsupervised machine learning that leverages recent advances in deep learning³ to compute approximate recursive equilibria in discrete-time models with (a) stochasticity, (b) a large amount of heterogeneity, which results in a high-dimensional state space, (c) financial frictions, which induce kinks in the equilibrium functions, and (d) an irregular geometry of the state space. The method we propose directly approximates the equilibrium functions using a deep neural network trained via a variant of mini-batch stochastic gradient descent on simulated data. We refer to our framework, which approximates all equilibrium functions directly, as *deep equilibrium nets* (DEQNs). The key idea of our proposed algorithm is to implement the loss function using the model's first-order conditions directly, thereby circumventing the need to obtain labeled data for supervised learning.⁴ The states used to train the DEQN are sampled by simulating the economy; by doing so, the network learns to approximate the equilibrium only where it matters: on an approximation of the ergodic set.⁵ In contrast to most grid-based methods, such as standard value function or policy function iteration with full Cartesian grids or sparse grids, our framework is able to approximate the equilibrium on irregularly shaped geometries. This property is one of the key elements for tackling high-dimensional models, as it avoids wasteful computations in parts of the state space where they are not needed (see, e.g., Maliar et al., 2011).

The main contribution of this article is that we introduce a grid-free, generic method based on simulations and deep neural networks to compute global solutions to high-dimensional and potentially highly nonlinear, dynamic stochastic models. Our method combines several desirable features. It does not rely on the invocation of nonlinear solvers or optimizers other than the gradient descent-based mechanism to update the parameters of the neural network. This allows for a swift generation of a substantial amount of training data via simulation. Moreover, through recent advances in neural network research that alleviate the curse of dimensionality,⁶ our method is capable of solving rich models that may be a better description of the real world than those used by practitioners to date. We also present a hybrid parallelization scheme based on Horovod (see Sergeev and Del Balso, 2018) that allows efficient use of contemporary high-performance computing hardware, which can speed up the training by orders of magnitude. To demonstrate the capabilities of our proposed algorithm, we solve three different models, all featuring a high-dimensional state space. That is, we solve two variations of

¹ For more details on Smolyak and adaptive sparse grids, see, for instance, Brumm and Scheidegger (2017), Judd et al. (2014), and Krueger and Kubler (2004).

² Adaptive sparse grids, for example, only address challenges (a), (b), and (c), whereas Gaussian processes (see Scheidegger and Bilionis, 2019) only address (a), (b), and (d).

³ We refer to Murphy (2012) for a general introduction to machine learning and to Goodfellow et al. (2016) for an introduction to deep learning.

⁴ Although this general notion is true for many applications of projection methods (see, e.g., Judd, 1992), our algorithm provides a concrete and simulation-based way to adapt this idea for the efficient use of deep learning.

⁵ The idea of simulating the economy on the ergodic set is not new. Haan and Marcet (1994), for instance, used simulation in the context of their parameterized expectation approach, which was later even extended to shallow neural networks by Duffy and McNelis (2001).

⁶ These include methodological advances (e.g., using rectified linear units (relu) as activation functions to facilitate backpropagation), improvements in the software (e.g., performance-optimized code libraries), and hardware (e.g., graphics and tensor processing units).

a classic overlapping generation (OLG) model from the literature, which we extend by adding borrowing constraints, adjustment costs, multiple assets, two types of agents, and an idiosyncratic health shock. Furthermore, we illustrate the broad applicability of our method by solving a Bewley (1977)-style model with a continuum of *ex ante* identical agents, per-period aggregate and idiosyncratic shocks, and recursive preferences. Code examples that illustrate our methodology are provided at <https://github.com/sischei/DeepEquilibriumNets>.

More specifically, we solve two annually calibrated life-cycle models with agents that live for 56 periods, borrowing constraints, multiple assets, and aggregate uncertainty. The second model, which we present in Appendix A.4, additionally features two types of agents and idiosyncratic health shocks. We chose this type of model for two reasons: first, this setup serves as a demonstrative example for the introduced method because the dimensionality of the state space increases with the number of periods the agents live for and, therefore, has a clear economic interpretation; second, because of its economic relevance. Explicitly accounting for an agent's life cycle is of crucial importance to, for instance, social security (see, e.g., Krueger and Kubler, 2006; Hasanhodzic and Kotlikoff, 2013), climate change (see, Kotlikoff et al., 2021a, 2021b, 2021c), labor (see, e.g., Gervais et al., 2016), and saving decisions. Saving decisions, for example, are driven by precautionary savings and life-cycle motives. The intergenerational wealth distribution hence affects consumption responses to a financial crisis or monetary policy (see, e.g., Wong, 2019). The rising life expectancy and quickly changing demographics render it a pressing task to understand how age and age distribution affect the economy. In our third application, we solve a Bewley (1977)-style model with a continuum of infinitely lived agents, aggregate and idiosyncratic risk, two regimes of aggregate uncertainty as well as Epstein–Zin preferences (Epstein and Zin, 1989, 1991). This application shows how our method can be applied to a model with a continuum of infinitely lived agents and recursive preferences.

Computationally, substantial challenges arise when working with life-cycle models in high-dimensional settings. In heterogeneous agent models with incomplete markets and uncertainty, traditional methods like value function iteration or time iteration (see, e.g., Judd, 1998; Ljungqvist and Sargent, 2000), in combination with Cartesian grid-based approximation schemes quickly become infeasible with increasingly many state variables. The curse of dimensionality (see Bellman, 1961)—that is, the exponential dependence of the overall computational effort on the number of dimensions—can be ameliorated by using Smolyak sparse grids (see Krueger and Kubler, 2004; Judd et al., 2014) or adaptive sparse grids (see Brumm and Scheidegger, 2017). However, a new challenge arises from these methods: the value functions or policy functions need to be approximated on a hypercube. Since the ergodic set of the states of the economy is frequently irregularly shaped, it often fills out only a tiny fraction of the hypercube (see also Judd et al., 2011). Thus, approximating the economy on a hypercubic domain can be wasteful and, in some cases, render the computation of solutions to models with irregularly shaped ergodic sets impossible. Furthermore, the researcher may not always be able to determine a hypercube in which the ergodic set of states of the economy lies. These problems can be addressed by grid-free simulation-based methods, which are suitable for high dimensions, such as Gaussian processes (see Rasmussen, 2004 for an introduction to Gaussian Processes and Scheidegger and Bilionis, 2019 for an application in economics). Gaussian processes, however, are not capable of dealing with a large amount of input data⁷ and hence cannot exploit the fact that simulation-based methods can generate input data for machine learning purposes at low cost. In contrast, neural networks turn out to successfully address all of these issues: they are suitable for dealing with very high-dimensional problems, they are a grid-free method, and they excel when there are much data⁸ available to learn from.

The algorithm introduced in this article can leverage these features of neural networks to approximate recursive equilibria in economic models with a high-dimensional state space. We

⁷ Standard Gaussian processes become computationally intractable for more than $\sim 10^4$ observations, as their computational complexity scales as $\mathcal{O}(N^3)$, where N is the number of observations.

⁸ That is, in the order of $\sim 10^6$ or more observations.

use the neural network to approximate all equilibrium functions directly. This allows us to simulate the economy at virtually zero cost, which in turn enables us to train the neural network on millions of points, which approximate the ergodic set in a grid-free fashion.

The remainder of the article is organized as follows: Section 2 gives a brief review of the related literature. Section 3 describes the benchmark model, which we use to illustrate our algorithm. Section 4 introduces the DEQN solution method. In Section 5, we showcase its performance in the context of the benchmark model. Section 6 concludes. In Appendix A.4, we show an additional application of DEQNs to an OLG model, similar to our benchmark model, but extended to feature two types of agents as well as an idiosyncratic health shock. In Appendix A.5, we further show how our method can be applied to solve a model with a continuum of infinitely lived agents with recursive preferences as well as aggregate and idiosyncratic risk. Appendix A.6 provides background information on neural networks more generally. In Appendix A.7, we discuss the remaining challenges and possible avenues for future research. In Appendix A.8, we present an application of DEQNs to a simple model, for which an analytical solution is known. Finally, Appendix A.9 describes our parallelization scheme.

2. LITERATURE REVIEW

The methodology we propose in this article contributes to three particular strands of literature on computing recursive equilibria numerically (see Figure A.1 in Appendix A.1 for an illustrated summary of the positioning of this article).

First, we propose a global solution method and thereby contribute to the literature focusing on the class of solution algorithms suitable for economic models featuring strong nonlinearities and a large amount of uncertainty. Second, our method is simulation-based and grid-free. Grid-free methods do not rely on constructing a rigid grid and, therefore, can approximate equilibria more efficiently⁹ on irregularly shaped state spaces. Finally, we contribute to the nascent strand of literature in computational economics that makes use of recent advances in machine learning to compute approximate equilibria in dynamic models. More precisely, we use unsupervised deep learning to approximate the equilibrium functions in annually calibrated life-cycle models directly.¹⁰ To the best of our knowledge, we are the first to do this. In the remainder of this section, we will outline the contributions of our algorithm and its relation to existing research in greater detail.

Since excellent reviews for the existing computational methods are available, our literature review focuses on placing our article within the emerging strand of literature that makes explicit use of machine learning to compute equilibria in economic models. Maliar and Maliar (2014), and Kollmann et al. (2011) give very detailed overviews of computational methods for economic models with a finite number of agents. For an overview of local and global methods, see Aruoba et al. (2006), Fernández-Villaverde et al. (2016), and Brumm et al. (2021).

The necessity of global solution methods for models with strong nonlinearities is pointed out, for instance, in Dou et al. (2017) and Fernández-Villaverde et al. (2015). (Smolyak) Sparse grids (see, e.g., Krueger and Kubler, 2004; Winschel and Krätsig, 2010; Judd et al., 2014) and adaptive sparse grids (see, e.g., Brumm et al., 2017; Brumm and Scheidegger, 2017) can ameliorate the curse of dimensionality but need to approximate the equilibrium on a hypercubic domain.¹¹ See, for example, Judd et al. (2011), Maliar and Maliar (2015), and Scheidegger and Bilionis (2019) for the advantages of grid-free, simulation-based methods for models with a high-dimensional state space and irregularly shaped ergodic sets.

⁹ Instead of, for example, interpolating the functions within a hypercube, where only a small fraction of the space that is interpolated may be relevant to the equilibrium, grid-free methods approximate the equilibrium functions at states that are reached during simulation.

¹⁰ With “directly”, we mean that the neural network approximates all policy functions in a way such that the economy can be simulated essentially by drawing the exogenous random shocks and by evaluating the neural network. This makes simulating the economy, a crucial part of our algorithm, fast.

¹¹ Brumm et al. (2017) solve a model similar to our benchmark model with a single asset. Using adaptive sparse grids to solve the model with two assets, as we do here, posed a prohibitive challenge for them.

The quest for computational methods, which can jointly address the obstacles of stochasticity, a high-dimensional state space, strong nonlinearities, and irregular state space geometries, led to the development of a new branch of grid-free methods, which makes explicit use of machine learning to compute equilibria in economic models. The highly active branch of machine learning research and the related literature from applied mathematics provide a rich set of powerful tools for approximating functions in high-dimensional settings, determining ergodic and feasible sets, and solving optimization problems efficiently—issues with which computational economic modeling is constantly confronted.

Neural networks are particularly promising since they have, to some extent, shown the potential to overcome the curse of dimensionality when solving partial differential equations and computing optimal stopping rules (Grohs et al., 2018; Jentzen et al., 2018; Sirignano and Spiliopoulos, 2018; Becker et al., 2019). Early applications of neural networks include Hutchinson et al. (1994), who hedge and price derivative securities, Chen and White (1999), who derive approximation rates for a particular type of time-series data, and Norets (2012), who uses neural networks in the context of estimation and discrete-state dynamic programming. Norets (2012), in particular, estimates finite-horizon, dynamic discrete choice models and uses a neural network to approximate expected value functions as a function of the economic parameters and state variables. Duffy and McNelis (2001) compare various methods for approximating and solving a stochastic growth model with parameterized expectations, one of which is a grid-free, simulation-based method that relies on shallow neural networks.

Closer to this article, Duarte (2018a, 2018b) presents a solution algorithm for economic models in continuous time, which uses neural networks to approximate the value function. The parameter updates, as implied by supervised learning, are computed efficiently by combining the Hamilton–Jacobi–Bellman equation with Ito's lemma and automatic differentiation. Fernández-Villaverde et al. (2019) use neural networks to forecast aggregate variables in a continuous-time model of financial frictions. Their algorithm builds on the seminal work of Krusell and Smith (1998) but uses neural networks to parameterize the perceived law of motion of aggregate variables, which allows the forecast of the aggregate variables to be nonlinear. In Fernández-Villaverde et al. (2019), the expectations of the households depend on a finite set of moments of the cross-sectional distribution of assets as well as an additional endogenous state variable (the expert's net wealth). The algorithm in Fernández-Villaverde et al. (2019) differs from ours along several dimensions. First, the neural network is only used to obtain the perceived law of motion, whereas the neural network in our method approximates all equilibrium functions. Second, the presented algorithm is developed for continuous-time models, whereas ours is developed for discrete time. Third, in contrast to Fernández-Villaverde et al. (2019), who, in addition, make use of equation solvers to compute the equilibrium, our algorithm uses simulation and a version of mini-batch stochastic gradient descent exclusively. In discrete-time settings, Villa and Valaitis (2019) use neural networks to approximate expectations in a grid-free manner to compute equilibria in a variety of settings where high-dimensional state spaces and the multicollinearity of state variables pose a computational challenge. Similar to our algorithm, their method iterates between a stochastic simulation phase and a learning phase. In contrast to Villa and Valaitis (2019), we do unsupervised learning.

In independent and contemporary work, Maliar et al. (2021) provide a general, grid-free framework for formulating economic models in a way that is suitable for the application of algorithms from the field of artificial intelligence. These formulations include the maximization of lifetime reward, the minimization of the Euler equation residuals, and the minimization of the Bellman residuals. As a test case, they solve the Krusell and Smith (1998) model with 2,000 state variables. Although the minimization of Euler equation residuals is similar to the algorithm proposed in this article, there are important differences. Next to the general idea of the algorithm, we consider it an equally important contribution to show when and how accurately it works. In contrast to Maliar et al. (2021), we focus on complex OLG models; we demonstrate the ability of our algorithm to solve them precisely by reporting comprehensive statistics on how well all equilibrium conditions are satisfied. Furthermore, we solve models

with two assets and collateral constraints, which is significantly more challenging computationally than comparable models with a single asset. In the application of our method to the model with a continuum of infinitely lived agents, we bring the Young (2010) approach to deep learning by including a histogram of asset holdings in the input to the neural network.¹² This is in contrast to Maliar et al. (2021), who encode the distribution as a collection of idiosyncratic asset holdings and idiosyncratic shocks of a large number of finitely many agents. In addition, we show how to embed recursive preferences into our algorithm.¹³

More recently, there are several papers making use of deep learning to solve challenging economic models. In continuous time, these include Fernández-Villaverde et al. (2020), who elegantly solve highly nonlinear models with up to 75 continuous state variables, as well as Gopalakrishna (2021), who incorporates economic constraints as regularizers and utilizes active learning to improve sampling of training points. In discrete time, Lepetyuk et al. (2020) use a deep learning framework to solve a challenging version of the “Terms of Trade Economic Model” of the Bank of Canada. Similarly, Maliar and Maliar (2022) solve a model with a single asset and discrete labor choice and Maliar and Maliar (2020) solve interesting neoclassical and New Keynesian single asset models with heterogeneous agents. Kahou et al. (2021) solve many agent models with neural networks and show that the exploitation of symmetries makes an essential contribution to breaking the curse of dimensionality. Fernández-Villaverde et al. (2021) apply the method developed by Fernández-Villaverde et al. (2019) in discrete time to study the interaction between the effect of the zero lower bound on aggregate quantities and household heterogeneity. In recent work, Folini et al. (2021) apply the method developed in this article to solve for the transition path in the context of climate economics. This can be done by extending the state space to include time. Han et al. (2021) introduce an efficient deep learning algorithm to solve heterogeneous agent models by approximating the distribution with a set of generalized moments.

3. THE BENCHMARK MODEL

In this section, we consider an OLG model with stochastic production very similar to that discussed in Krueger and Kubler (2004). In this model, the dimensionality of the state space increases linearly with the number of age groups. Hence, a high-dimensional state space arises naturally and in an economically meaningful way. To demonstrate the applicability of our method in a setting with multiple assets and occasionally binding constraints, we extend the model of Krueger and Kubler (2004) along two lines: first, we include borrowing constraints and adjustment costs on capital, which renders it an illiquid asset; second, we allow agents to trade a liquid one-period bond subject to collateral constraints. In addition to the computational challenges, these extensions, therefore, allow the model to connect to literature highlighting the role of assets with different liquidity on the cross-sectional consumption response to aggregate shocks (see, e.g., Kaplan et al., 2018; Wong, 2019).

3.1. Uncertainty. The OLG model we consider is in discrete time—that is, $t = 0, 1, \dots$. In each period, one of Z possible discrete exogenous shocks realizes. We assume that the shocks follow a first-order Markov process with finite support $\mathcal{Z} := \{1, \dots, Z\}$ with transition matrix Π . We denote the aggregate shock realizing in period t by z_t and a history of aggregate shocks up to period t by $z^t = (z_0, \dots, z_t)$.

3.2. Households. We study an OLG model where agents live for N periods. There is no uncertainty about the lifetime, and there is one representative household per cohort. At each

¹² The nonstochastic simulation approach for idiosyncratic shocks has two advantages: first, the histogram encodes the distribution of asset holdings directly, which is economically meaningful information and may simplify training; second, nonstochastic simulation circumvents the noisiness of Monte Carlo simulations.

¹³ Recursive preferences are a particularly important use case for our algorithm since recursive preferences are commonly used in asset pricing, where global solution methods are often required.

node z^t , a representative household is born. Households only distinguish themselves by their birth-node $z^{t^{\text{birth}}}$. At time $t \in \{t^{\text{birth}}, \dots, t^{\text{birth}} + N - 1\}$, we identify the household born at t^{birth} by its current age $s = t - t^{\text{birth}} + 1$. For example, the consumption of the household with age s at period t is denoted by $c^s(z^t)$. We omit the explicit dependence on z^t where there is no risk of confusion and write c_t^s . In each period, the alive agents receive a strictly positive labor endowment, which only depends on the age of the agent. The labor endowment of an agent of age s is denoted by l^s . The price of the consumption good is normalized to one. Furthermore, we assume that households supply their labor endowment inelastically for a market wage $w(z^t) = w_t$. At each node z^t , the alive agents maximize their remaining time-separable discounted expected lifetime utility given by

$$(1) \quad \sum_{i=0}^{N-s} E_t [\beta^i u(c_{t+i}^{s+i})],$$

where the discount factor $\beta < 1$. Furthermore, we assume that the utility function $u : \mathbb{R}_{++} \rightarrow \mathbb{R}$ is smooth, strictly increasing, strictly concave, and that it satisfies the Inada condition $\lim_{c \rightarrow 0} u'(c) = \infty$.

There are two ways for households to transfer consumption over time. First, households can save in risky capital, which is subject to adjustment costs, and hence illiquid. The illiquid savings of household s in period t are denoted by $a^s(z^t) = a_t^s$. The savings will become capital in the next period:

$$(2) \quad a_t^s = k_{t+1}^{s+1}, \quad \forall t, \quad \forall s \in \{1, \dots, N - 1\},$$

where k_t^s denotes illiquid capital holdings of age group s in the beginning of period t . Households sell their capital to the firm at market price r_t . The return to capital accrues into the household's illiquid account. The deposit into the illiquid account by household s in period t is given by

$$(3) \quad \Delta_t^s = a_t^s - k_t^s r_t.$$

Capital is illiquid due to the convex adjustment costs given by

$$(4) \quad \frac{\zeta}{2} (\Delta_t^s)^2,$$

where ζ is the intensity of the capital adjustment costs. Furthermore, we impose an exogenous borrowing limit \underline{a} , which can be set to zero when borrowing is prohibited:

$$(5) \quad a_t^s \geq \underline{a}.$$

Next to saving in illiquid, risky capital, households can buy a one-period bond at market price p_t . The bond is in zero net supply. A bond promises a payoff of 1 in the subsequent period. Let b_t^s denote the bond holding of household s in period t and let d_t^s denote the amount of bonds purchased by household s in period t , so that

$$(6) \quad b_{t+1}^{s+1} = d_t^s.$$

The bond is liquid in the sense that there are no costs associated with adjusting the bond holding. However, selling the bond is subject to collateral constraints. To model collateral constraints, we follow Kubler and Schmedders (2003) and use

$$(7) \quad \kappa d_t^s + a_t^s \geq 0,$$

where κ is an exogenous constant, and only capital serves as collateral. If a bond-selling household defaults on the payment in the following period, the value of the corresponding amount of capital is transferred to the bond-buying households; otherwise, there is no punishment.¹⁴ Consequently, households only default if the value of the collateral is smaller than the promised payout of the bond. The realized payout of the bond is then given by

$$(8) \quad \min\{\kappa r_t, 1\}.$$

In our numerical experiments below, we chose κ high enough so that the bond is indeed a risk-free asset, and default does not occur in equilibrium. Therefore, $\min\{\kappa r_t, 1\} = 1$, which we use in the following for clearer notation. The budget constraint of household s in period t is therefore given by

$$(9) \quad c_t^s + p_t d_t^s + a_t^s + \frac{\zeta}{2} (\Delta_t^s)^2 = l^s w_t + b_t^s + r_t k_t^s.$$

Finally, the agents are born and die without any assets—that is, $k_t^1 = b_t^1 = 0$, and $a_t^N = d_t^N = 0$.

3.3. Firms. There is a single representative firm with a Cobb–Douglas production function, where the total factor productivity (TFP) and the depreciation depend on the exogenous shock alone. Each period, after the shock has been realized, the firm buys capital and hires labor to maximize its profits, taking prices as given. The production function is given by

$$(10) \quad f(K_t, L_t, z_t) = \eta_t K_t^\alpha L_t^{1-\alpha} + K_t(1 - \delta_t),$$

where K_t denotes the aggregate capital bought, L_t denotes the aggregate labor hired, α denotes the capital share in production, η_t denotes the stochastic TFP, and δ_t denotes the stochastic depreciation.

3.4. Markets. The price of capital (r_t) and wages (w_t), as well as the price of the bond (p_t), are determined by market clearing in competitive spot markets for consumption, capital, labor, and bonds.

3.5. Equilibrium. Following Krueger and Kubler (2004), and accounting for our extensions to the model, we now define a competitive equilibrium for our economy:

DEFINITION 1 (COMPETITIVE EQUILIBRIUM). A competitive equilibrium, given initial conditions $z_0, \{k_0^s\}_{s=1}^N, \{b_0^s\}_{s=1}^N$, is a collection of choices for households $\{(c_t^s, a_t^s, d_t^s)\}_{s=1}^N$ and for the representative firm $(K_t, L_t)_{t=0}^\infty$ as well as prices $(r_t, w_t, p_t)_{t=0}^\infty$, such that

1. Given $(r_t, w_t)_{t=0}^\infty$, the choices $\{(c_t^s, a_t^s, d_t^s)\}_{s=1}^N$ maximize (1), subject to (2), (5), (6), (7), and (9).
2. Given r_t, w_t , the firm maximizes profits:

$$(11) \quad (K_t, L_t) \in \arg \max_{K_t, L_t \geq 0} f(K_t, L_t, z_t) - r_t K_t - w_t L_t.$$

3. All markets clear: For all t ,

$$(12) \quad L_t = \sum_{s=1}^N l_t^s, \quad K_t = \sum_{s=1}^N k_t^s, \quad 0 = \sum_{s=1}^N b_t^s.$$

¹⁴ To be precise, we assume that in case of default, the value of the pledged collateral, κr_t , is transferred between the liquid accounts.

For our choice of the production function, given in Equation (10), the first-order conditions of the firm's maximization problem imply that

$$(13) \quad w_t = (1 - \alpha)\eta_t K_t^\alpha L_t^{-\alpha},$$

$$(14) \quad r_t = \alpha\eta_t K_t^{\alpha-1} L_t^{1-\alpha} + (1 - \delta_t).$$

For future reference, we write down the household's optimality conditions as well. The Karush–Kuhn–Tucker (KKT) conditions for any given generation of age $s \in 1, \dots, N-1$ at node z^t are given by

$$(15) \quad (1 + \zeta \Delta_t^s) u'(c_t^s) = \beta E_t[u'(c_{t+1}^{s+1})r_{t+1}(1 + \zeta \Delta_{t+1}^{s+1})] + \lambda_t^s + \mu_t^s,$$

$$(16) \quad \lambda_t^s \cdot (a_t^s - \underline{a}) = 0,$$

$$(17) \quad a_t^s - \underline{a} \geq 0,$$

$$(18) \quad \lambda_t^s \geq 0,$$

$$(19) \quad p_t u'(c_t^s) = \beta E_t[u'(c_{t+1}^{s+1})] + \kappa \mu_t^s,$$

$$(20) \quad \mu_t^s \cdot (a_t^s + \kappa d_t^s) = 0,$$

$$(21) \quad a_t^s + \kappa d_t^s \geq 0,$$

$$(22) \quad \mu_t^s \geq 0.$$

The generation of terminal age N simply consumes everything it has.

4. APPROXIMATION OF EQUILIBRIA WITH DEEP NEURAL NETWORKS

To describe how to solve the OLG model presented in Section 3 with DEQNs, we proceed in two steps. In Subsection 4.1, we define a functional rational expectations equilibrium (FREE) for the presented economy. Subsection 4.2 describes how to use deep neural networks to search for an approximate recursive equilibrium by using a projection method (see, e.g., Judd, 1992 and Gaspar and Judd, 1997). In Appendix A.9, we additionally provide a generic parallelization scheme that can speed up our methodology by orders of magnitude.

4.1. Functional Rational Expectations Equilibrium. Our proposed algorithm aims at approximating a recursive equilibrium: a function mapping the state of the economy to choices and prices, which are consistent with the equilibrium conditions. Following Spear (1988) and Krueger and Kubler (2004), we call such a function a *functional rational expectations equilibrium*. For our presented economy, the state of the economy can be described by the current exogenous shock and the distribution of assets holdings across age groups. More precisely, we define a FREE as follows:

DEFINITION 2 (FUNCTIONAL RATIONAL EXPECTATIONS EQUILIBRIUM). A FREE consists of equilibrium functions $\boldsymbol{\theta} = [\boldsymbol{\theta}_a^T, \boldsymbol{\theta}_\lambda^T, \boldsymbol{\theta}_d^T, \boldsymbol{\theta}_\mu^T, \theta_p]^T : \mathcal{Z} \times \mathbb{R}^{2N} \rightarrow \mathbb{R}^{4(N-1)+1}$, where $\boldsymbol{\theta}_a : \mathcal{Z} \times \mathbb{R}^{2N} \rightarrow \mathbb{R}^{N-1}$ denotes the capital investment functions, $\boldsymbol{\theta}_\lambda : \mathcal{Z} \times \mathbb{R}^{2N} \rightarrow \mathbb{R}^{N-1}$ denotes the KKT-multiplier functions for the short-selling constraint on capital, $\boldsymbol{\theta}_d : \mathcal{Z} \times \mathbb{R}^{2N} \rightarrow \mathbb{R}^{N-1}$ denotes the bond investment functions, $\boldsymbol{\theta}_\mu : \mathcal{Z} \times \mathbb{R}^{2N} \rightarrow \mathbb{R}^{N-1}$ denotes the KKT-multiplier functions for the collateral constraint, and $\theta_p : \mathcal{Z} \times \mathbb{R}^{2N} \rightarrow \mathbb{R}$ denotes the bond price function, such that for all states $\mathbf{x} := [z, \mathbf{k}^T, \mathbf{b}^T]^T \in \mathcal{Z} \times \mathbb{R}^{2N}$, where $z \in \mathcal{Z}$ denotes the exogenous shock, and the capital holding $\mathbf{k} = [k_1, \dots, k_N]^T$ together with the bond holding $\mathbf{b} = [b_1, \dots, b_N]^T$ denote the endogenous state (i.e., the distribution of capital holdings and bond holdings) with $k_1 = 0$ and

$b_1 = 0$, we have for all $i = 1, \dots, N - 1$:¹⁵

$$(23) \quad (1 + \zeta \Delta(\mathbf{x})_i) u'(c(\mathbf{x})_i) = \beta E_z[(r(\mathbf{x}_+)(1 + \zeta \Delta(\mathbf{x}_+)_{i+1}) u'(c(\mathbf{x}_+)_{i+1})) + \theta_\lambda(\mathbf{x})_i + \theta_\mu(\mathbf{x})_i],$$

$$(24) \quad 0 = \theta_\lambda(\mathbf{x})_i \theta_a(\mathbf{x})_i,$$

$$(25) \quad 0 \leq \theta_a(\mathbf{x})_i,$$

$$(26) \quad 0 \leq \theta_\lambda(\mathbf{x})_i,$$

as well as

$$(27) \quad \theta_p(\mathbf{x}) u'(c(\mathbf{x})_i) = \beta E_z[u'(c(\mathbf{x}_+)_{i+1})] + \kappa \theta_\mu(\mathbf{x})_i,$$

$$(28) \quad 0 = \theta_\mu(\mathbf{x})_i (\theta_a(\mathbf{x})_i + \kappa \theta_d(\mathbf{x})_i),$$

$$(29) \quad 0 \leq \theta_a(\mathbf{x})_i + \kappa \theta_d(\mathbf{x})_i,$$

$$(30) \quad 0 \leq \theta_\mu(\mathbf{x})_i,$$

and

$$(31) \quad 0 = \sum_{i=1}^{N-1} \theta_d(\mathbf{x})_i,$$

where

$$(32) \quad \mathbf{x}_+ = [z_+, 0, \boldsymbol{\theta}_a(\mathbf{x})^T, 0, \boldsymbol{\theta}_d(\mathbf{x})^T]^T,$$

where z_+ denotes the random exogenous shock in the next period, and where

$$(33) \quad \Delta(\mathbf{x})_i := \theta_a(\mathbf{x})_i - r(\mathbf{x})x_{1+i},$$

$$(34) \quad r(\mathbf{x}) = f_K \left(\sum_{i=1}^N x_{1+i}, \sum_{i=1}^N l^i(x_1), x_1 \right),$$

$$(35) \quad w(\mathbf{x}) = f_L \left(\sum_{i=1}^N x_{1+i}, \sum_{i=1}^N l^i(x_1), x_1 \right),$$

$$(36) \quad c(\mathbf{x})_i = l^i w(\mathbf{x}) + x_{1+i+N} - \theta_p(\mathbf{x}) \theta_d(\mathbf{x})_i - \Delta(\mathbf{x})_i - \frac{\zeta}{2} (\Delta(\mathbf{x})_i)^2.$$

Computing an approximate FREE in this model translates to finding an approximation for the $4 \cdot (N - 1) + 1$ equilibrium functions such that the above equations are approximately fulfilled for all exogenous shocks and all values the $2N$ -dimensional endogenous state takes.

4.2. Algorithm to Train DEQNs. The goal of our solution framework is to approximate the equilibrium functions $\boldsymbol{\theta}$ (see Definition 2) with a deep neural network. To do so, we combine four components. Generically, the four components are given by: (i) a suitable class of function approximators; (ii) a loss function measuring the quality of a given approximation at a given state; (iii) an updating mechanism to improve the approximation; and (iv) a sampling method for choosing states for the updating and the evaluation of the approximation quality.

¹⁵ With a slight abuse of notation, we set $\theta_a(\mathbf{x}_+)_N = \theta_d(\mathbf{x}_+)_N = 0$. More precisely, $\theta_a(\mathbf{x}_+)$ and $\theta_d(\mathbf{x}_+)$ are $N - 1$ -dimensional vectors. To be formally correct, we would need to distinguish between $i < N - 1$ and $i = N - 1$ when writing down the equations.

Specific to our algorithm, we chose deep neural networks as a function approximator. The loss function is implemented using the errors in the equilibrium conditions and the neural network parameters are updated using variants of mini-batch gradient descent. To update the parameters of the neural network, as well as to evaluate the quality of our approximation, we sample states from a simulated path of the economy.

This combination of the four components allows us to (a) use a large number of states to assess the quality of our approximation, (b) sample the states from the approximated ergodic distribution of states in the economy, (c) handle irregular geometries of the ergodic set of states, and (d) approximate equilibrium functions with kinks and strong nonlinearities. We next outline each of the components separately and then combine them into one algorithm.

4.2.1. Function approximator. We use densely connected feedforward neural networks as function approximators because they combine a set of desirable qualities. Neural networks are universal function approximators (Hornik et al., 1989), can resolve distinct local, highly nonlinear features, and can handle a large amount of high-dimensional input data.¹⁶

Given hyperparameters $\{K, \{m_i\}_{i=1}^K, \{\sigma_i(\cdot)\}_{i=1}^K\}$ and trainable parameters ρ , a neural network \mathcal{N}_ρ encodes the map

$$(37) \quad \mathbf{x} \rightarrow \mathcal{N}_\rho(\mathbf{x}) = \sigma_K(\mathbf{W}_K \dots \sigma_2(\mathbf{W}_2 \sigma_1(\mathbf{W}_1 \mathbf{x} + \mathbf{b}_1) + \mathbf{b}_2) \dots + \mathbf{b}_K),$$

where $\mathbf{W}_i \in \mathbb{R}^{m_{i+1} \times m_i}$ are matrices often referred to as weight matrices, and $\mathbf{b}_i \in \mathbb{R}^{m_{i+1}}$ are vectors often referred to as bias vectors. The vector ρ denotes the collection of all entries of the weight matrices and the bias vectors. K is referred to as the number of layers of the neural network and m_i as the number of nodes in layer i . The nonlinear functions σ_i are referred to as activation functions and are applied elementwise to each entry of a vector: $\sigma_i(\mathbf{x}) = [\sigma_i(x_1), \dots, \sigma_i(x_{m_{i+1}})]^T$. A densely connected feedforward neural network is hence given by a sequence of matrix-vector multiplications followed by the application of an activation function.

4.2.2. Loss function. The goal of our algorithm is to approximate the equilibrium functions θ (see Definition 2) by a neural network. In this section, we will introduce a loss function: a measure of the quality of our approximation at a given state of the economy.

Let ρ denote the set of trainable parameters of the neural network and let the neural network, given the set of parameters ρ , be denoted by \mathcal{N}_ρ . The neural network maps the state \mathbf{x} into the approximated equilibrium functions

$$(38) \quad \mathcal{N}_\rho : \mathcal{Z} \times \mathbb{R}^{2N} \rightarrow \mathbb{R}^{4(N-1)+1} : \mathbf{x} \rightarrow \mathcal{N}_\rho(\mathbf{x}),$$

where $\mathbf{x} = [z, \mathbf{k}^T, \mathbf{b}^T]^T$, and

$$(39) \quad \begin{aligned} \mathcal{N}_\rho(\mathbf{x}) &= \hat{\theta}(\mathbf{x}) = [\hat{\theta}_a(\mathbf{x})^T, \hat{\theta}_\lambda(\mathbf{x})^T, \hat{\theta}_d(\mathbf{x})^T, \hat{\theta}_\mu(\mathbf{x})^T, \hat{\theta}_p(\mathbf{x})^T] \\ &=: [\hat{a}(\mathbf{x})_1, \dots, \hat{a}(\mathbf{x})_{N-1}, \hat{\lambda}(\mathbf{x})_1, \dots, \hat{\lambda}(\mathbf{x})_{N-1}, \hat{d}(\mathbf{x})_1, \dots, \hat{d}(\mathbf{x})_{N-1}, \hat{\mu}(\mathbf{x})_1, \dots, \hat{\mu}(\mathbf{x})_{N-1}, \hat{p}(\mathbf{x})]^T. \end{aligned}$$

Since the true equilibrium functions θ are defined by the conditions given in Definition 2, our aim is to find parameters ρ , such that the resulting approximation of the policy functions fulfills Equations (23)–(36) for all $i = 1, \dots, N - 1$. Given neural network parameters ρ and a state \mathbf{x}_j of the economy, we define a measure of the quality of the approximation by evaluating the errors in the equilibrium conditions, which result from the policy functions encoded by the neural network.

¹⁶ See Appendix A.6 for more background information on neural networks and Goodfellow et al. (2016) for a more comprehensive introduction to deep learning.

For readability, we will define the error in each of the equations separately before summing them to form a single measure encoding all equilibrium conditions.

Firm optimization. Some of the equilibrium conditions can be enforced to hold exactly for all states. For any state \mathbf{x} and neural network parameters ρ , the equilibrium prices for capital and labor depend on the state alone. We can determine the correct equilibrium prices for capital and labor using Equations (34) and (35) and, therefore, satisfy them exactly.

Budget constraint of the household. We can also ensure that the budget constraints of the households hold exactly by substituting $\hat{\mathbf{a}}$, $\hat{\mathbf{d}}$, and \hat{p} , which are encoded by neural network parameters ρ , into the equilibrium policies in Equation (36):

$$(40) \quad \hat{c}(\mathbf{x})_i = l^i w(\mathbf{x}) + x_{1+i+N} - \hat{p}(\mathbf{x}) \hat{d}(\mathbf{x})_i - \hat{\Delta}(\mathbf{x})_i - \frac{\zeta}{2} (\hat{\Delta}(\mathbf{x})_i)^2,$$

where

$$(41) \quad \hat{\Delta}(\mathbf{x})_i := \hat{a}(\mathbf{x})_i - r(\mathbf{x}) x_{1+i}.$$

State transition. The transition from one state to the next, Equation (32), can also be satisfied exactly by setting the next period's endogenous state consistent with the current period's policies:

$$(42) \quad \mathbf{x}_+ = [z_+, 0, \hat{\mathbf{a}}(\mathbf{x})^T, 0, \hat{\mathbf{d}}(\mathbf{x})^T]^T.$$

Inequalities. There are four inequalities (Equations (25), (26), (29), and (30)) from the remaining equilibrium conditions which can be enforced through the architecture of the neural network.

Specifically, using an activation function that has a nonnegative range in the output layer, such as the softplus function, can ensure that the inequalities are always satisfied. To allow for this, we redefine the neural network's output such that it approximates $\theta_{\text{col}}(\mathbf{x}) := \theta_a(\mathbf{x}) + \kappa \theta_d(\mathbf{x})$, where $\theta_{\text{col}}(\mathbf{x})$ is the collateral requirement policy, instead of approximating $\theta_d(\mathbf{x})$ directly.¹⁷ That is,

$$\mathcal{N}_\rho(\mathbf{x}) = [\hat{\theta}_a(\mathbf{x})^T, \hat{\theta}_\lambda(\mathbf{x})^T, \hat{\theta}_{\text{col}}(\mathbf{x})^T, \hat{\theta}_\mu(\mathbf{x})^T, \hat{\theta}_p(\mathbf{x})^T]^T.$$

Household's optimization. The equilibrium conditions related to the household's optimization problem are Equations (23), (24), (27), and (28). As opposed to the previously discussed conditions, we cannot easily enforce them for all possible states. Therefore, taking the enforced equilibrium conditions, the neural network parameters ρ , and a given state \mathbf{x}_j as given, we construct a measure for how well the current approximation fulfills each of the remaining conditions.

Following Judd (1992), we convert the Euler equation error of generation i in state \mathbf{x}_j (corresponding to Equation (23)) to the relative Euler equation error implied by the neural network policy function defined as

$$(43) \quad e_{\mathbf{x}_j}^{i, \text{REE cap}}(\rho) := \frac{u'^{-1} \left(\frac{\beta E_{z_j} [r(\mathbf{x}_{j,+})(1+\zeta \hat{\Delta}(\mathbf{x}_{j,+})_{i+1}) u'(\hat{c}(\mathbf{x}_{j,+})_{i+1})] + \hat{\lambda}(\mathbf{x}_j)_i + \hat{\mu}(\mathbf{x}_j)_i}{1+\zeta \hat{\Delta}(\mathbf{x}_j)_i} \right)}{\hat{c}(\mathbf{x}_j)_i} - 1.$$

¹⁷ The approximation of the bond policy is easily backed out from the neural network by $\hat{\theta}_d(\mathbf{x}) = (\hat{\theta}_{\text{col}}(\mathbf{x}) - \hat{\theta}_a(\mathbf{x}))/\kappa$.

The advantage of using the relative Euler equation error is that its value has an economic interpretation that is independent of the utility function; namely, it quantifies the consumption error. Evaluating the Euler equation requires the evaluation of the expectation operator. In the models presented in this article, we work with discrete shocks and can therefore evaluate the expectation operator exactly. In case of continuous shocks, the expectation operator could be approximated with standard numerical integration methods.¹⁸

Similarly, we define the relative Euler equation error for bond investment (corresponding to Equation (27)) as

$$(44) \quad e_{\mathbf{x}_j}^{i, \text{REE bond}}(\rho) := \frac{u'^{-1}\left(\frac{\beta E_{z_j}[u'(\hat{c}(\mathbf{x}_{j+})_{i+1}] + \kappa \hat{u}(\mathbf{x}_j)_i}{\hat{\rho}(\mathbf{x}_j)}\right)}{\hat{c}(\mathbf{x}_j)_i} - 1.$$

The errors in the remaining KKT conditions (Equations (24) and (28)) are given by

$$(45) \quad e_{\mathbf{x}_j}^{i, \text{KKT cap}}(\rho) := \hat{\lambda}(\mathbf{x}_j)_i \hat{a}(\mathbf{x}_j)_i, \text{ and}$$

$$(46) \quad e_{\mathbf{x}_j}^{i, \text{KKT bond}}(\rho) := \hat{\mu}(\mathbf{x}_j)_i (\hat{a}(\mathbf{x}_j)_i + \kappa \hat{d}(\mathbf{x}_j)_i).$$

Market clearing. The final equilibrium condition, Equation (31), is the market clearing condition of the bond. We define the error in the market clearing condition as

$$(47) \quad e_{\mathbf{x}_j}^{\text{mc}}(\rho) = \sum_{i=1}^{N-1} \hat{d}(\mathbf{x}_j)_i.$$

Loss function encoding the equilibrium conditions. If the neural network \mathcal{N}_ρ would encode the true equilibrium functions exactly, Equations (43)–(47) would evaluate to zero for all states \mathbf{x} of the economy. Therefore, we can define a loss function, that is, a measure of the quality of the approximation \mathcal{N}_ρ , by quantifying the extent to which Equations (43)–(47) differ from zero when evaluated at a given state \mathbf{x} . Our unsupervised loss function generally depends on two components: the parameters of the function approximator ρ and the set of states \mathbf{x} at which the desired conditions are evaluated. The latter is referred to as the *training set*, which we denote as $\mathcal{D}_{\text{train}}$. Given parameters ρ and a set of states $\mathcal{D}_{\text{train}}$, we define the loss function as the mean squared error of all equilibrium conditions:¹⁹

$$(48) \quad \ell_{\mathcal{D}_{\text{train}}}(\rho) := \frac{1}{|\mathcal{D}_{\text{train}}|} \sum_{\mathbf{x}_j \in \mathcal{D}_{\text{train}}} \left(\frac{1}{N-1} \sum_{i=1}^{N-1} \left(e_{\mathbf{x}_j}^{i, \text{REE cap}}(\rho) \right)^2 + \left(e_{\mathbf{x}_j}^{i, \text{REE bond}}(\rho) \right)^2 + \left(e_{\mathbf{x}_j}^{i, \text{KKT cap}}(\rho) \right)^2 + \left(e_{\mathbf{x}_j}^{i, \text{KKT bond}}(\rho) \right)^2 + \left(e_{\mathbf{x}_j}^{\text{mc}}(\rho) \right)^2 \right).$$

4.2.3. Updating mechanism. This subsection describes how to use the loss function to optimize the trainable parameters ρ .²⁰ The loss function is defined such that smaller values

¹⁸ The computational cost of approximating the expectation operator grows linearly with the number of quadrature nodes used.

¹⁹ If, for reasons implied by the economic model, particular equilibrium conditions require higher precision than others, the relative weighting of the different conditions can be adjusted. Ultimately, it is essential that the loss function, which the neural network optimizes, is aligned with the implied loss function the researcher aims to optimize.

²⁰ The hyperparameters have to be chosen by the modeler prior to the training we describe in this section. The modeler may use prior experience, manual, random, or grid search as well as methods such as Bayesian optimization (see, e.g., Bergstra et al., 2011).

correspond to lower mean squared errors in the equilibrium conditions. Consequently, parameters are deemed “good” if they minimize the loss function. Due to the functional structure of deep neural networks, variants of gradient descent are typically used to optimize the parameters ρ .²¹ Gradient descent updates the parameters stepwise in the direction in which the loss function decreases—that is:

$$(49) \quad \rho_k^{\text{new}} = \rho_k^{\text{old}} - \alpha^{\text{learn}} \frac{\partial \ell_{\mathcal{D}_{\text{train}}}(\rho^{\text{old}})}{\partial \rho_k^{\text{old}}} \quad \forall k \in \{1, \dots, \text{length}(\rho)\}.$$

The parameter $\alpha^{\text{learn}} > 0$ that governs by how much the parameters are adjusted with each gradient descent step is referred to as the *learning rate*. Popular variants of gradient descent that can speed up the learning process are briefly discussed in Appendix A.6.3. Deep and wide neural networks that are trained on limited training data $\mathcal{D}_{\text{train}}$ tend to overfit. That is, the neural network optimizes the parameters ρ on the training set $\mathcal{D}_{\text{train}}$ to the extent that the approximation quality deteriorates for new, unseen states, not part of the training set. Hence, the availability of training data is often a bottleneck in deep learning applications. The formulation of our loss function together with an efficient algorithm to generate new and large training sets $\mathcal{D}_{\text{train}}$ (as introduced in the next section) enables our algorithm to circumvent this bottleneck.

4.2.4. Sampling. As described in Subsection 4.2.3, the parameters of the neural network ρ are chosen to minimize the loss function (Equation (48)). In this section, we describe the training set ($\mathcal{D}_{\text{train}}$) generation process used to obtain a good approximation of the equilibrium functions for the ergodic set of states of the economy.

Since we want to use the approximated equilibrium functions to simulate the modeled economy, they must provide a good approximation for those states, which will be visited during the simulation. Therefore, we chose to train the neural network on the states visited on the simulated path of the economy. To do so, we start with an arbitrary, economically feasible starting state \mathbf{x}_1^0 , and randomly initialized neural network parameters ρ . Then, we simulate $T - 1$ periods forward based on the approximated equilibrium functions given by the neural network.²² Since our method approximates the equilibrium functions directly, simulating the evolution of the economy is computationally cheap. The resulting T simulated states of the economy constitute our data set $\mathcal{D}_{\text{train}}^0 = \{\mathbf{x}_1^0, \dots, \mathbf{x}_T^0\}$. We call the set of T simulated periods $\mathcal{D}_{\text{train}}$ an *episode*. We split this input data into mini-batches—smaller subsets of size m with random membership—and perform gradient descent steps on each subset as given in Equation (49). A completion of an *epoch* is defined as when the whole data set $\mathcal{D}_{\text{train}}$ is passed through the algorithm. Per epoch, the neural network parameters are updated T/m times. Next, we set $\mathbf{x}_1^1 = \mathbf{x}_T^0$ and use the updated parameters of the neural network to simulate $T - 1$ periods forward, generate a new training set $\mathcal{D}_{\text{train}}^1 = \{\mathbf{x}_1^1, \dots, \mathbf{x}_T^1\}$, and repeat the process. Since we can generate a large amount of training data in this setting, overfitting is not a primary concern.²³ As the neural network learns better parameter values, the simulated states become better representatives of the ergodic set of states of the economy.

Since the loss function merely requires a set of feasible states of the economy to be evaluated, our algorithm does not require us to obtain the training data $\mathcal{D}_{\text{train}}$ from simulations. If the ergodic set of states of the economy is known, if one wants to approximate the equilibrium functions on a larger set or if one wants to improve the approximation quality at specific areas of the state space, the training set $\mathcal{D}_{\text{train}}$ can be chosen accordingly.

²¹ We use Adam (see Kingma and Ba, 2014) together with mini-batch gradient descent. See Appendix A.6.3 for details.

²² Since the neural network parameters are initialized randomly, some corrections have to be made for the case when the neural network predicts an infeasible endogenous state in the next period (such as negative aggregate capital). The adjustments are easy to make and are described in Appendix A.7.3.

²³ Our method allows us to train the neural network on more than a billion simulated states.

4.2.5. Algorithm. In this section, we summarize how the components described in the previous sections are combined to compute approximate recursive equilibria with DEQNs.

Starting from a randomly initialized neural network, our algorithm iterates between generating a new training set $\mathcal{D}_{\text{train}}$ by simulating a desired amount of states and improving the parameters ρ of the neural network by performing a variant of gradient descent steps on the training set. The error on a new set of simulated states is an out-of-sample error and can be used to judge the out-of-sample quality of the approximation. Because we use the neural network to approximate the equilibrium functions directly, the simulation of a new set of points is computationally cheap. Therefore, we can set the number of epochs to train on each set of simulated points to one. Consequently, each simulated state is only used for a single gradient descent step, which guards our algorithm against overfitting. Algorithm 1 provides the pseudo-code of the DEQN.²⁴ Our parallelization scheme, which can speed up the computations by orders of magnitude, is outlined in Appendix A.9.

Algorithm 1. Algorithm for training deep equilibrium nets

Data:
 T (length of an episode), N_{epochs} (number of epochs on each episode), N_{iter} (maximum number of iterations),
 $\tau^{\text{mean}}, \tau^{\text{max}}$ (desired threshold for mean and max error, respectively),
 $\epsilon^{\text{mean}} = \infty, \epsilon^{\text{max}} = \infty$ (starting value for current mean and max error, respectively),
 ρ^0 (initial parameters of the neural network), x_1^0 (initial state to start simulations from), $i = 0$ (set counter),
 α^{learn} (learning rate)

Result:
success (boolean if thresholds were reached)
 ρ^{final} (final neural network parameters)

```

while (( $i < N_{\text{iter}}$ )  $\wedge$  ( $(\epsilon^{\text{mean}} \geq \tau^{\text{mean}}) \vee (\epsilon^{\text{max}} \geq \tau^{\text{max}})$ )) do
     $\mathcal{D}_{\text{train}}^i \leftarrow \{x_1^i, x_2^i, \dots, x_T^i\}$  (generate new training data)
     $x_0^{i+1} \leftarrow x_T^i$  (set new starting point)
     $\epsilon_{\text{max}} \leftarrow \max \left\{ \max_{x \in \mathcal{D}_{\text{train}}^i} |e_x \cdot (\rho)| \right\}, \epsilon_{\text{mean}} \leftarrow \max \left\{ \frac{1}{T} \sum_{x \in \mathcal{D}_{\text{train}}^i} |e_x \cdot (\rho)| \right\}$  (update errors)
    for  $j \in [1, \dots, N_{\text{epochs}}]$  do
        (learn  $N_{\text{epochs}}$  on data)
        for  $k \in [1, \dots, \text{length}(\rho)]$  do
            (50)  $\rho_k^{i+1} = \rho_k^i - \alpha^{\text{learn}} \frac{\partial \ell_{\mathcal{D}_{\text{train}}^i}(\rho^i)}{\partial \rho_k^i}$ 
            (do a gradient descent step to update the network parameters)
        end
    end
     $i \leftarrow i + 1$  (update episode counter)
end
if  $i = N_{\text{iter}}$  then return (success  $\leftarrow \text{False}, \rho^{\text{final}} \leftarrow \rho^i$ );
else return (success  $\leftarrow \text{True}, \rho^{\text{final}} \leftarrow \rho^i$ );
```

5. DEQNS IN ACTION: SOLVING THE BENCHMARK MODEL

To evaluate the performance of our algorithm, we study an annually calibrated OLG model, as outlined in Section 3. We chose this model because it features the computational challenges we want to address with our solution framework. We model $N = 56$ age groups—that is, the endogenous state is 110-dimensional.²⁵ Furthermore, there are aggregate shocks, and occasionally binding constraints. In consequence, the OLG model specification used here

²⁴ For simplicity, the pseudo-code implements gradient descent instead of mini-batch gradient descent.

²⁵ To be precise, a sufficient endogenous state is 108-dimensional since agents enter the economy without any assets.

offers a unique setting to illustrate the performance of our new solution method for an economically relevant class of models.

In addition, in Appendix A.4, we solve a more complicated model with two sources of idiosyncratic risk over the life cycle. The first source of idiosyncratic risk is being born into one of two trajectories of labor endowment over the life cycle. The second source of idiosyncratic risk is a potential health shock shortly before retiring. The state space in this model is 292-dimensional. Nevertheless, we demonstrate that our method solves the model accurately. We use this stylized model to study the welfare benefits of health insurance, modeled as an asset that pays out contingent on the realization of the health shock.

In the following, we first summarize the parameterization of the benchmark model and the hyperparameters of the neural network. We then demonstrate that our proposed algorithm can solve the model with high accuracy. In Appendix A.3, we study the aggregate and cross-sectional consumption response to aggregate shocks.

5.1. Parameterization. In this section, we briefly outline the model parameters (cf., Subsection 5.1.1) and neural network hyperparameter specification (cf., Subsection 5.1.2) that we use in our numerical experiments below.

5.1.1. Model parameters. A summary of our parameterization is given in Table 1. We keep the parameterization standard. Details are given in Appendix A.2. Importantly, we have four exogenous aggregate shocks: two depreciation shocks and two shocks to TFP, which evolve independently of each other. Shocks 1 and 2 correspond to normal times with a capital depreciation of $\delta = 0.08$. Shocks 3 and 4 are high depreciation states with a depreciation of $\delta = 0.11$, which is roughly in line with the literature on rare disasters (see, e.g., Gourio, 2012; Usui, 2019). In both, normal and disaster states, TFP takes values in $\eta \in \{0.978, 1.022\}$ with a probability of 0.905 to remain in the same state. We model $N = 56$ age groups, corresponding to ages 25–80. The parameterization of the age-dependent labor endowment is based on Guvenen et al. (2021) and Brumm et al. (2017) (see Appendix A.2 for details).

5.1.2. Neural network hyperparameters. We use a deep neural network with two hidden layers to solve our benchmark model. Heuristically, we found it to be helpful to augment the state of the economy with redundant information before passing it to the neural network.²⁶ The input layer consists of $12 + 4 \cdot N$ input nodes, which for $N = 56$ results in 236 input nodes. After the input layer, the neural network features two hidden layers with 1,000 relu-activated hidden nodes each. The output layer consists of $4 \cdot (N - 1) + 1 = 221$ nodes, activated with softplus functions to ensure that the nonnegativity constraints are fulfilled.

A schematic illustration of the neural network architecture, for clarity displayed without providing redundant information, is given in Figure 1.²⁷

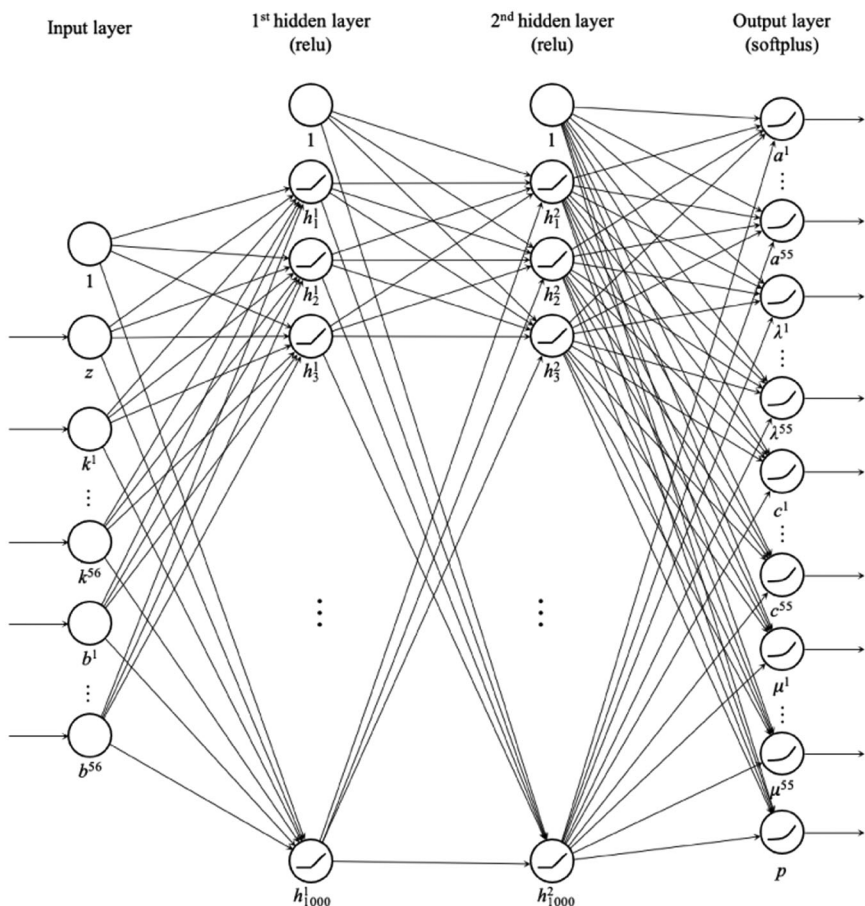
We keep the number of epochs per episode N^{epochs} small to avoid overfitting. The learning rate has to be chosen small enough, and the mini-batch size large enough, so that the mini-batch gradient descent steps are not too noisy. Heuristically, we found it helpful to decrease the learning rate and increase the mini-batch size toward the end of the training procedure in order to fine tune the neural network parameters. Therefore, we switch from mini-batch size $m = 64$ and learning rate of $\alpha^{\text{learn}} = 10^{-5}$ to a larger mini-batch size $m = 1,000$ and a smaller

²⁶ See Appendix A.7.2 for more details.

²⁷ As typical when working neural networks, the architecture presented here is not the result of theoretically motivated performance optimization, but based on numerical experimentation. We choose the activation function of the output layer to encode economic constraints (here nonnegativity) into the neural network architecture. For the activation functions of the hidden layers, there is little theoretical guidance on which activation functions to choose. Relu activations are the de facto standard for densely connected neural networks as they mitigate the vanishing gradient problem (in contrast to sigmoid and TanH) and promote sparsity in connections. Our numerical experiments did not provide evidence that other hidden layer activation functions improve performance.

TABLE 1
PARAMETERIZATION OF THE BENCHMARK MODEL (CF, SECTION 3)

		Persistence		Depreciation		Collateral Requirement	
		Persistence TFP	Depreciation	$P(\delta_{t+1} = 0.08 \delta_t = 0.08)$	Borrowing Constraint	Adjustment Cost	Capital ζ
Discount Factor β	Relative Risk Aversion γ	Capital Share α	TFP η	Depreciation δ	$P(\delta_{t+1} = 0.11 \delta_t = 0.11)$	\underline{a}	κ
0.95	2	0.3	[0.978, 1.022]	[0.08, 0.11]	0.905	0.972	0.5
					0.700	0.0	1.1236



NOTES: Schematic illustration of the neural network architecture for the deep equilibrium net. For simplicity, it is illustrated for the case where no redundant information is passed to the neural network. See Appendix A.7 for explanations on providing additional inputs to the neural network.

FIGURE 1

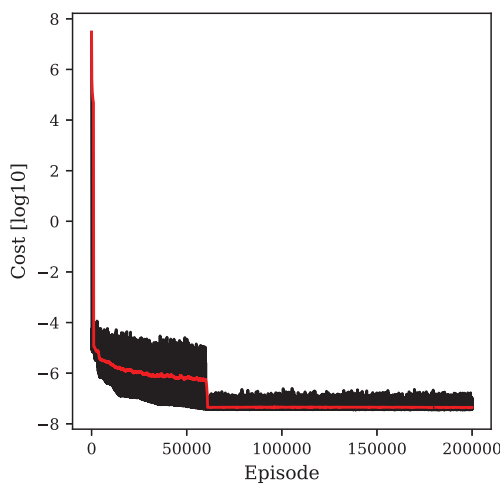
SCHEMATIC ILLUSTRATION OF THE NEURAL NETWORK ARCHITECTURE

TABLE 2
HYPERPARAMETERS CHOSEN TO TRAIN THE DEEP EQUILIBRIUM NET FOR THE BENCHMARK MODEL

Episodes	Learning Rate α^{learn}	Periods per Episode T	Epochs per Episode N_{epochs}	Mini-Batch Size m	Nodes Hidden Layers	Activations Hidden Layers
1–60,000	1×10^{-5}	10,000	1	64	1,000	relu
					1,000	relu
					1,000	relu
60,000–200,000	1×10^{-6}	10,000	1	1,000	1,000	relu

learning rate $\alpha^{\text{learn}} = 10^{-6}$ after 60,000 episodes of training.²⁸ The chosen hyperparameters are specified in Table 2. To accelerate the simulation, we simulate 1,000 10-period paths in parallel, instead of a single 10,000-period path.

²⁸ Increasing the batch size has the effect that the gradient estimates are less noisy. Decreasing the learning rate results in smaller steps being taken. The initial hyperparameters are higher to accelerate the initial rough-grain stage of training.



NOTES: Evolution of the cost function during 200,000 episodes of training, 1 epoch per episode. The black line shows the cost function evaluated on a newly simulated episode, the red line shows a moving average over the last 1,000 episodes. The corresponding neural network parameters are described in Subsection 5.1.

FIGURE 2

EVOLUTION OF THE COST FUNCTION

TABLE 3
STATISTICS OF ERRORS IN THE EQUILIBRIUM CONDITIONS

	Mean	Max	0.1	10	50	90	99.9
Rel Ee capital [%]	0.024	0.528	0.000	0.002	0.013	0.060	0.324
Rel Ee bond [%]	0.021	0.608	0.000	0.002	0.012	0.051	0.255
KKT capital [$\times 10^{-2}$]	0.000	0.127	0.000	0.000	0.000	0.000	0.000
KKT bond [$\times 10^{-2}$]	0.005	0.104	0.000	0.000	0.000	0.000	0.021
Market clearing [$\times 10^{-2}$]	0.053	0.333	0.000	0.017	0.047	0.089	0.252
Market clearing (normalized) [$\times 10^{-2}$]	0.000	0.002	0.000	0.000	0.000	0.000	0.001

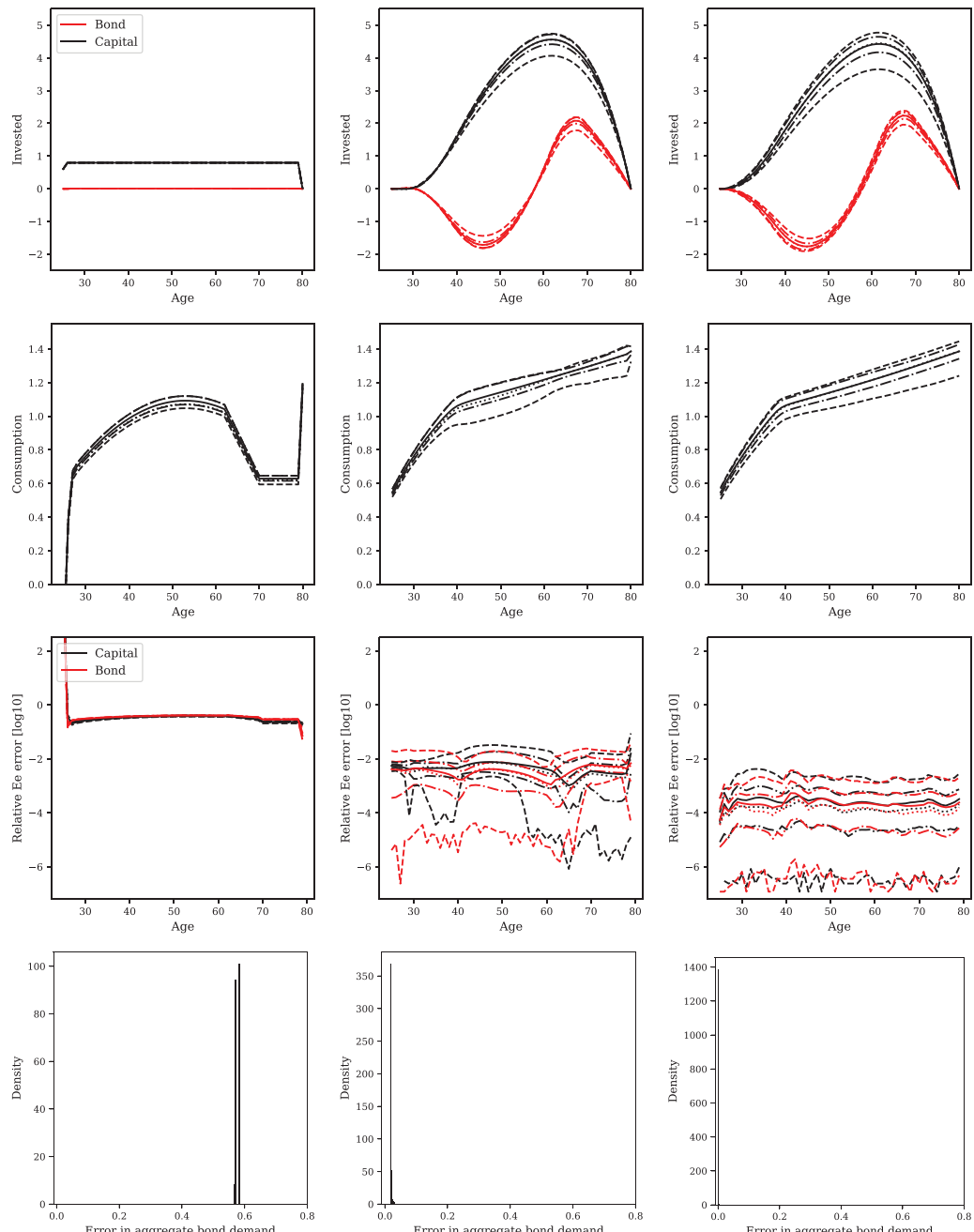
NOTE: The first two rows show the relative Euler equation errors (Rel Ee in %) on 10,000 simulated periods. Rows 3 and 4 show the errors in the KKT conditions ($\times 10^{-2}$). Rows 5 and 6 show the errors in the market clearing conditions ($\times 10^{-2}$). In row 6, the error in the market clearing condition for the bond is divided by aggregate production. The columns show the mean and the max errors as well as the 0.1st, 10th, 50th, 90th, and 99.9th percentile.

5.2. Training Progress of the DEQN. Next, we illustrate the performance of the proposed algorithm for our benchmark model. Figure 2 shows the evolution of the cost function during the first 200,000 episodes of training: the cost function decreases quickly during training and converges below $10^{-7.0}$. The step-like decrease after 60,000 episodes stems from the increase in the mini-batch size and the decrease in the learning rate, as described in the previous section.²⁹

Figure 3 shows the distribution of investments in capital and the bond (first row), consumption (second row), and relative Euler equation errors (third row), as well as error in the bond market clearing conditions (fourth row) on a simulated path of 2,000 simulated states after 1, 1,000, and 200,000 episodes of training. As the training process advances, it can be seen that the DEQN learns better approximations of the policies (first and second rows) and that the errors in the equilibrium conditions decrease (third and fourth rows).

Table 3 reports the mean and maximum relative Euler equation error (in %), error in the KKT conditions ($\times 10^{-2}$), and market clearing error ($\times 10^{-2}$), as well as

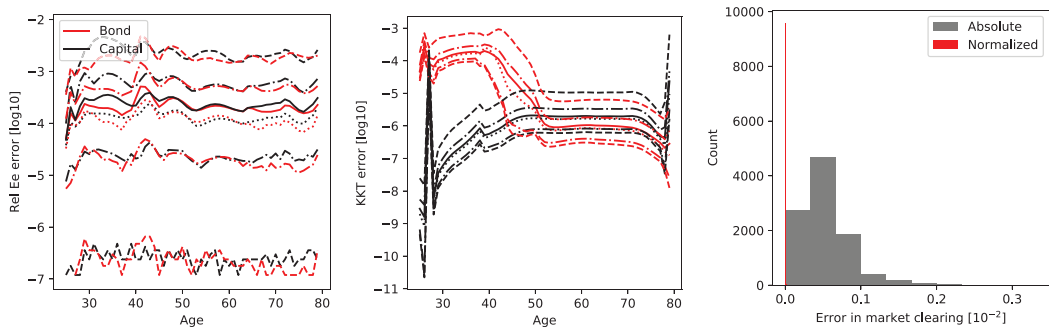
²⁹ The training of the benchmark model typically consumes less than 10 minutes of compute time on a single hybrid CPU/GPU node to reach mean relative Euler equation errors below 1%.



NOTES: Distribution of investment (first row), consumption (second row), relative Euler equation errors (third row), and errors in the market clearing condition for the bond on a simulated path after learning for 1 (left column), 1,000 (middle column), and 200,000 (right column) episodes. The solid line shows the mean over 2,000 simulated periods, the dotted line shows the median, the dash-dotted lines show the 10th and 90th percentile, the dashed lines show the 0.1st and 99.9th percentile.

FIGURE 3

EVOLUTION OF POLICIES AND ERRORS DURING TRAINING



NOTES: Distribution of the relative Euler equation error (left), error in the KKT conditions (middle), and errors in the market clearing condition (right). The solid line shows the mean over 10,000 simulated periods, the dotted line shows the median, the dash-dotted lines show the 10th and 90th percentile, the dashed lines show the 0.1st and 99.9th percentile.

FIGURE 4

ERRORS AFTER TRAINING

several percentiles on a simulated path of 10^4 periods, after training the DEQN for 2×10^5 episodes.³⁰

The mean relative error for all equilibrium conditions is below 0.1%. A relative Euler equation error of 0.1% means that the agents, on average, consume 0.1% too much or too little. The 99.9th percentile and the maximum of all errors is below 1%. The left and center panels in Figure 4 plot the errors across the age groups and the right panel plots the errors in the market clearing condition.

The reached accuracy demonstrates that our method can compute satisfactory approximations to the FREE.

In Appendix A.3, we use the computed equilibrium functions to study the aggregate and distributional consumption response to aggregate shocks. An application of our method to an extended life-cycle model, which additionally features two types of agents and an idiosyncratic health shock, is presented in Appendix A.4. Appendix A.5 presents a further application of our method to a model with a continuum of agents, aggregate, and idiosyncratic shocks.

6. CONCLUSION

This article introduces DEQNs—neural networks that approximate all equilibrium functions and are trained on simulated data to satisfy all equilibrium conditions. Since the neural network approximates the equilibrium functions directly, neither sets of nonlinear equations nor optimization problems need to be solved to simulate the economy. Consequently, training data can be generated at virtually zero cost, which allow us to swiftly train the neural network on more than a billion simulated states of the economy. DEQNs thus provide a grid-free, global solution method, which can jointly address the computational challenges arising from a high-dimensional state space, significant uncertainty, financial frictions, and irregular geometries of the state space. The training process is fully parallelizable, which allows efficient use of contemporary high-performance computing hardware.

We illustrate the performance of our method in the context of three different models. We solve two OLG models with a 108 and 292-dimensional endogenous state space, occasionally binding constraints, and aggregate uncertainty. Both models feature two types of assets: risky, illiquid capital and a risk-free, one-period bond subject to collateral requirements. In Appendix A.3, we study the aggregate and cross-sectional consumption response

³⁰ We simulate a total of 12,000 periods and discard the first 2,000 “burn-in” periods (see, e.g., Krusell and Smith, 1998).

to aggregate shocks in the benchmark model and find significant heterogeneity across age groups. The second model, which we present in Appendix A.4, additionally features two types of agents and an idiosyncratic health shock. In Appendix A.5, we further show how our method naturally extends to models with a continuum of infinitely lived agents with recursive utility in the presence of aggregate and idiosyncratic risk. We bring the Young (2010) method to deep learning and solve a Bewley (1977)-style model, extended with aggregate risk, two regimes of aggregate uncertainty, as well as Epstein–Zin preferences.

All examples show that our proposed method can accurately approximate recursive equilibria in economically relevant settings with a high-dimensional state space, uncertainty, and strong nonlinearities.

ACKNOWLEDGMENTS

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APPENDIX

A.1 Supplementary Figures.

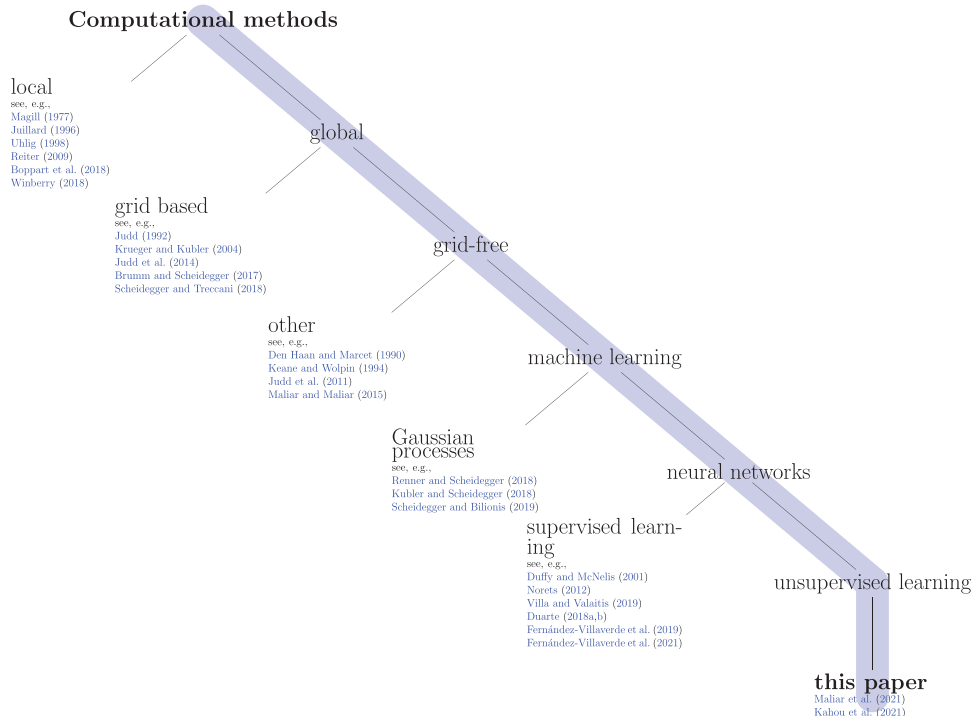


FIGURE A.1

ILLUSTRATION OF THE PLACEMENT OF THIS PAPER WITHIN THE LITERATURE (NONEXHAUSTIVE LIST)

A.2 Details on the Benchmark Model Parameters. We study an overlapping generation (OLG) model when borrowing in capital is prohibited entirely, that is, $\underline{a} = 0$.

The stochastic depreciation in our model aims to represent rare disasters in a parsimonious and stylized way. In the spirit of Gourio (2012), we chose a depreciation of $\delta = 0.08$ during normal times and $\delta = 0.11$ in the disaster state. The probability to enter a disaster during normal times is given by 0.028 and the persistence of the disaster state is chosen to be

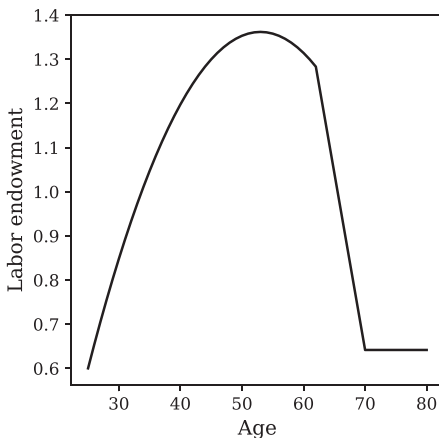


FIGURE A.2

LABOR ENDOWMENT OVER THE LIFE CYCLE IN THE BENCHMARK MODEL

0.70, resulting in an average disaster length of roughly 3.33 years. The associated transition matrix is given by

$$(A.1) \quad \Pi^\delta = \begin{bmatrix} 0.972 & 0.028 \\ 0.300 & 0.700 \end{bmatrix},$$

which is in line with values, which are used in the literature on rare disasters and capital depreciation (see, e.g., Gourio, 2012; Usui, 2019).

The total factor productivity (TFP) process is modeled as an independent AR(1) process, with an unconditional average of $\bar{\eta} = 1$, a yearly persistence of $\rho_\eta = 0.81$, and normally distributed yearly innovations with mean zero and standard deviation of $\sigma_\eta = 0.013$. These values are within the standard range of the real business cycle literature (see, e.g., Fernández-Villaverde and Guerron-Quintana, 2020). We discretize the AR(1) process into a two-state Markov process following Rouwenhorst (1995). More precisely, TFP η takes one of two values, $\eta \in \{0.978, 1.022\}$ and evolves according to the transition matrix

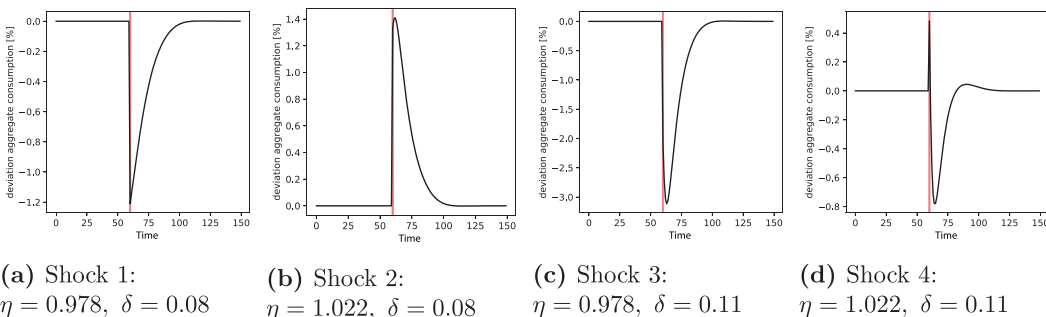
$$(A.2) \quad \Pi^\eta = \begin{bmatrix} 0.905 & 0.095 \\ 0.095 & 0.905 \end{bmatrix}.$$

Depreciation and TFP evolve independently of each other. In total, there are four possible combinations of depreciation shock and TFP shock that are labeled by $z \in \mathcal{Z} = \{1, 2, 3, 4\}$ and correspond to $\delta = [0.08, 0.08, 0.11, 0.11]^T$ and $\eta = [0.978, 1.022, 0.978, 1.022]^T$. The overall transition matrix is given by $\Pi = \Pi^\delta \otimes \Pi^\eta$, where $\Pi_{i,j}$ denotes the probability of a transition from the exogenous shock i to exogenous shock j , where $i, j \in \mathcal{Z}$.

The labor endowment over the life cycle is shown in Figure A.2.

Agents are “born” with age 25, work until age 63, and are retired from ages 64 to 80. During the agents’ working age, the labor endowment is hump shaped and increases by 127% between age 25 to its peak value at age 53. The shape and rise of the labor endowment roughly mimics the average earnings by age group documented by Guvenen et al. (2021). Following Brumm et al. (2017), we model the retirement period by a linearly decreasing segment, followed by a constant segment. That is, the labor endowment drops by 50% between ages 64 and 70, whereafter it stays constant.

Following Krueger and Kubler (2004), the capital share of production is set to $\alpha = 0.3$, patience is set to $\beta = 0.95$, and the agents’ utility from consumption exhibit constant relative risk



NOTES: Mean percent deviation of the aggregate consumption from its unconditional average when each of the exogenous shocks occurs in period 60. The mean impulse response was computed over 100,000 simulations. On impact of the respective shock, aggregate consumption changes as follows: (a) decreases by 1.2%; (b) increases by 1.3%; (c) decreases by 2.1%; and (d) increases by 0.5%, respectively.

FIGURE A.3

IMPULSE RESPONSE OF AGGREGATE CONSUMPTION

aversion of $\gamma = 2$. The parameter κ , which determines the collateral requirement, is set, such that agents can borrow at most as much as is covered by the minimum value of their depreciated capital in the next period: $\kappa = \frac{1}{1-0.11} \approx 1.12$. The bond is hence a risk-free asset. The parameter ζ , governing the adjustment costs of capital, is set to 0.5. See Table 1 for a summary of the economic model parameters.

A.3 Consumption Response to Aggregate Shocks in the Benchmark Model. We now use the equilibrium functions that we computed in Section 5.2 to study the consumption response to aggregate shocks across age groups. We thereby relate to recent literature, which highlights the importance of the cross-sectional consumption response and the liquidity of assets for understanding the aggregate consumption response to economic shocks (in the context of monetary policy, see, e.g., Kaplan et al., 2018; Wong, 2019). First, we analyze the impulse response of aggregate consumption to each of the four shocks. Next, we show how the consumption response to aggregate shocks varies across age groups.

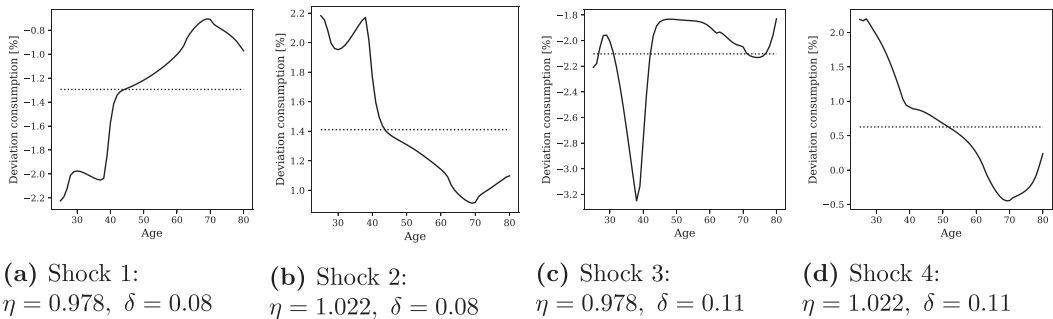
To study the reaction of consumption to each of the four aggregate shocks, we perform the following exercise. We draw 10^5 starting states from the ergodic distribution.³¹ From each starting state, we simulate 150 periods forward twice. In one simulation, the exogenous shock evolves randomly for all 150 periods. In the other, we specify that a particular shock realizes in the period 60. For each time period and each case, with and without a specified shock in the period 60, we compute averages of the quantities of interest. Then, we compare the time series of average economic quantities.

First, we study the impulse response of aggregate consumption to each of the four shocks, which is shown in Figure A.3. On average, aggregate consumption is about 1.2% lower when the low TFP shock realizes together with normal depreciation and about 1.3% higher when high TFP realizes. In the event of the rare high depreciation shocks, aggregate consumption drops by 2.1% when it realizes together with low TFP and, on impact, increases by 0.5% when it realizes together with high TFP.³²

Next, we want to understand the response of aggregate consumption by looking at the response across different age groups. Figure A.4 shows the cross-sectional reaction of consumption to each of the four shocks in the period in which the shock realizes. The dotted lines show the mean consumption deviation, averaged across age groups.

³¹ We simulate 1,000 different stochastic evolutions of the economy with 50,000 periods each, and pick simulated states 500 periods apart as our starting states.

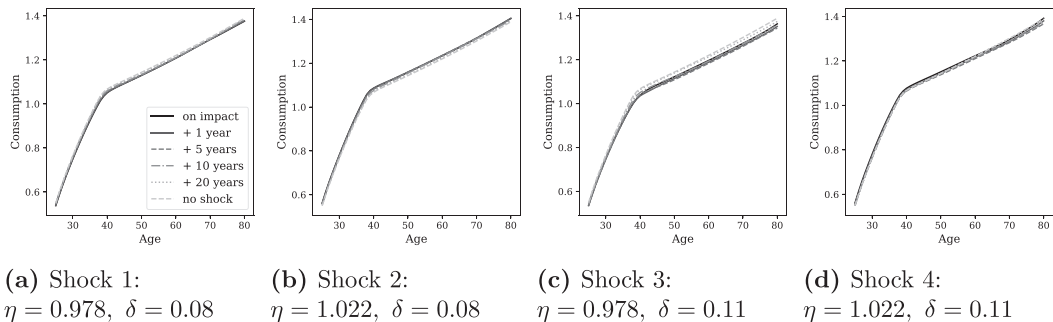
³² Note that in the latter case, the consumption response is positive on impact, but then negative in the period after.



NOTES: Mean percent deviation of consumption by age group from its unconditional average when each of the exogenous shocks occurs. The dotted lines show the mean across age groups. The mean is taken over 100,000 simulations.

FIGURE A.4

CONSUMPTION RESPONSE ACROSS AGE GROUPS



NOTES: The mean is taken over 100,000 simulations.

FIGURE A.5

MEAN CONSUMPTION ACROSS AGE GROUPS AFTER REALIZATION OF AN AGGREGATE SHOCK

As the figure shows, there is significant heterogeneity across age groups, where young agents respond the strongest. When the high depreciation shock realizes together with low TFP, these agents earn a low return on their illiquid asset and become borrowing constrained and hence have to reduce their debt, resulting in lower consumption. Agents between 40 and 50 hold a larger level of debt, however they hold enough illiquid wealth so that the collateral constraint rarely binds.

To see the impact of the aggregate shock on cross-sectional consumption in the subsequent periods, Figure A.5 shows the average consumption across age groups in subsequent periods. The impact of experiencing a high depreciation shock on consumption remains visible for more than 20 years.

A.4 A Model with Two Types of Agents, Overlapping Generations, and Idiosyncratic Health Shocks. The benchmark model presented in Section 3 featured one type of agent and no idiosyncratic risk. Next, we want to enrich the model by adding two sources of idiosyncratic risk over the life cycle. First, we now consider two possible trajectories of labor endowment over the life cycle, reflecting two types of agents in the economy. The type of an agent is persistent and realizes at "birth." Second, agents face a health shock shortly before retiring. Health risks are an important source of uncertainty over the life cycle, especially in our aging society, and subject to the public and academic debate (see, e.g., De Nardi et al., 2016; Pelgrin and St-Amour, 2016). Hence, we believe that questions related to health insurance are naturally linked to the intergenerational wealth distribution, rendering OLG models a

suitable modeling choice. This endorses the need for appropriate solution methods, such as the method we present in this article. In the stylized model, we present in this section, agents experience a reduced utility from consumption in the event of falling sick, causing them to draw heavily on their savings, and hence leaving them a significantly reduced amount of savings when they enter retirement. In this setting, we then study the welfare gains from introducing health insurance, that is, the possibility to invest in an additional asset, which pays out contingent on the realization of the health shock. This section hence serves to illustrate that our method is perfectly applicable in the presence of several idiosyncratic shocks as well as to richer models, which may help address policy-relevant questions.

A.4.1 The economic model. Except for the two types of agents and the health shocks, the model remains identical to our benchmark model. Therefore, we introduce the new model with a focus on the differences with respect to the benchmark model, which is described in Section 3.

An agent in this economy is indexed by the tuple (y, s, h) , and $y \in \{1, 2\}$ denotes the two types of agents. In every cohort, there is a mass one of agents of each type. The agent's age is indexed by $s \in \{1, \dots, N\}$, and $h \in \{1, 2\}$ denotes whether the agent experienced a health shock ($h = 2$) or not ($h = 1$). The index h evolves according to a type-specific Markov transition probability $\Pi^{(y,s)}$. We assume that the health shock realizes at fixed age $s = s^h$.³³ The fraction of agents who receive the health shock at age s^h is given by $\pi^y := \Pi_{1,2}^{(y,s^h-1)}$. Consequently, we have

$$(A.3) \quad \Pi^{(y,s)} = \mathbb{1}_{s < s^h-1} \begin{bmatrix} 1 & 0 \\ 0 & 0 \end{bmatrix} + \mathbb{1}_{s=s^h-1} \begin{bmatrix} 1 - \pi^y & \pi^y \\ 0 & 0 \end{bmatrix} + \mathbb{1}_{s \geq s^h} \begin{bmatrix} 1 & 0 \\ 0 & 1 \end{bmatrix}.$$

We chose s^h to be the last period before retirement. The per-period utility function in this model is given by

$$(A.4) \quad u\left(\frac{c_t^{(y,s,h)}}{g^{(y,s,h)}}\right),$$

where the factor $g^{(y,s,h)} \geq 1$ models the health shock. If an agent of type y and age $s = s^h$ experiences the health shock, that is, $h = 2$, the agent draws less utility of consumption by a factor $g^{(y,s^h,2)} > 1$. For all other agents, we have $g^{(y,s,h)} = 1$, and the utility from consumption is unchanged. For $s < s^h$, h only takes value 1 for both types. For $s \geq s^h$, we have two agents per type and cohort corresponding to $h = 1$ and $h = 2$.

The rest of the model follows our benchmark model, which is described in Section 3. We denote the agents' holdings in risky, illiquid capital by $k_t^{(y,s,h)}$, the investment is denoted by $a_t^{(y,s,h)}$, where $a_t^{(y,s,h_t)} = k_{t+1}^{(y,s+1,h_{t+1})}$. Similarly, $b_t^{(y,s,h)}$ and $d_t^{(y,s,h)}$ denote bond holdings and bond purchases, respectively, where $d_t^{(y,s,h_t)} = b_{t+1}^{(y,s+1,h_{t+1})}$. The borrowing constraints on capital and the collateral requirement for short-selling the bond remain also as in the benchmark model.

In our benchmark model, there is a single representative agent per age group, resulting in N heterogeneous agents that interact every period. In contrast, we now have $\tilde{N} = (s^h - 1) + 2 \cdot (N - (s^h - 1))$ agents for each type y . That is, there is one agent per period younger than s^h and two agents per period from age s^h onward, where the members of the latter cohort distinguish themselves through whether they experienced the health shock. Therefore, for two types $y \in \{1, 2\}$, we model a total of $2 \cdot \tilde{N}$ agents. In our model calibration, $N = 56$ and $s^h = 38$

³³ Since the health shock only occurs at age s^h , the transition of index h is deterministic for all age groups other than for age group $s = s^h - 1$.

(corresponding to age 62 when agents are born at 25), resulting in 150 agents and an approximately 300-dimensional endogenous state.³⁴

A.4.2 Functional rational expectations equilibrium (FREE). In this section, we define a FREE for the OLG model with idiosyncratic shocks. Note that next to a larger number of agents, the only equilibrium conditions which change are the households' optimality conditions for age groups $s = s^h - 1$ (i.e., in the period before the health shock realizes), and $s = s^h$ (i.e., in the period, in which the health shock realizes). We denote the \tilde{N} policy functions of type y for investment in capital with $a^{(y,s,h)}(\cdot)$, with associated Karush–Kuhn–Tucker (KKT) multipliers $\lambda^{(y,s,h)}(\cdot)$, and the policy functions for investment in the bond with $d^{(y,s,h)}(\cdot)$, with associated KKT multipliers $\mu^{(y,s,h)}(\cdot)$. The price function for the bond is denoted by $p(\cdot)$.

DEFINITION A.1 (FUNCTIONAL RATIONAL EXPECTATIONS EQUILIBRIUM). A FREE consists of equilibrium functions $\{a^{(y,s,h)}(\cdot), \lambda^{(y,s,h)}(\cdot), d^{(y,s,h)}(\cdot), \mu^{(y,s,h)}(\cdot), p(\cdot)\}$, where $a^{(y,s,h)} : \mathcal{Z} \times \mathbb{R}^{2\tilde{N}} \rightarrow \mathbb{R}^{\tilde{N}-2}$ denotes the capital investment functions, $\lambda^{(y,s,h)} : \mathcal{Z} \times \mathbb{R}^{2\tilde{N}} \rightarrow \mathbb{R}^{\tilde{N}-2}$ denotes the KKT-multiplier functions for the short-selling constraint on capital, $d^{(y,s,h)} : \mathcal{Z} \times \mathbb{R}^{2\tilde{N}} \rightarrow \mathbb{R}^{\tilde{N}-2}$ denotes the bond investment functions, and $\mu^{(y,s,h)} : \mathcal{Z} \times \mathbb{R}^{2\tilde{N}} \rightarrow \mathbb{R}^{\tilde{N}-2}$ denotes the KKT-multiplier functions for the collateral constraint for type $y \in \{1, 2\}$, such that for all states $\mathbf{x} := [z, \mathbf{k}^1, \mathbf{k}^2, \mathbf{b}^1, \mathbf{b}^2]^T \in \mathcal{Z} \times \mathbb{R}^{2\tilde{N}}$, where $z \in \mathcal{Z}$ denotes the exogenous shock, and the capital holding \mathbf{k}^y together with the bond holding \mathbf{b}^y denote the endogenous state (i.e., the distribution of capital holdings and bond holdings) with $k^{(y,1,1)} = 0$ and $b^{(y,1,1)} = 0$, we have for all $s = 1, \dots, N-1, y = 1, 2, h = 1$ for $s < s^h$ and $h = 1, 2$ for $s \geq s^h$:

$$\begin{aligned} & \frac{1 + \zeta \Delta^{(y,s,h)}(\mathbf{x})}{g^{(y,s,h)}} u' \left(\frac{c^{(y,s,h)}(\mathbf{x})}{g^{(y,s,h)}} \right) \\ &= \beta \mathbb{E}_{z,h} \left[\frac{r(\mathbf{x}_+) (1 + \zeta \Delta^{(y,s+1,h_+)}(\mathbf{x}_+))}{g^{(y,s+1,h_+)}} u' \left(\frac{c^{(y,s+1,h_+)}(\mathbf{x}_+)}{g^{(y,s+1,h_+)}} \right) \right] \\ & \quad + \lambda^{(y,s,h)}(\mathbf{x}) + \mu^{(y,s,h)}(\mathbf{x}), \end{aligned} \tag{A.5}$$

$$0 = a^{(y,s,h)}(\mathbf{x}) \cdot \lambda^{(y,s,h)}(\mathbf{x}), \tag{A.6}$$

$$0 \leq a^{(y,s,h)}(\mathbf{x}), \tag{A.7}$$

$$0 \leq \lambda^{(y,s,h)}(\mathbf{x}), \tag{A.8}$$

as well as

$$\begin{aligned} & \frac{p(\mathbf{x})}{g^{(y,s,h)}} u' \left(\frac{c^{(y,s,h)}(\mathbf{x})}{g^{(y,s,h)}} \right) \\ &= \beta \mathbb{E}_{z,h} \left[\frac{1}{g^{(y,s+1,h_+)}} u' \left(\frac{c^{(y,s+1,h_+)}(\mathbf{x}_+)}{g^{(y,s+1,h_+)}} \right) \right] \\ & \quad + \kappa \mu^{(y,s,h)}(\mathbf{x}), \end{aligned} \tag{A.9}$$

$$0 = \mu^{(y,s,h)}(\mathbf{x}) (a^{(y,s,h)}(\mathbf{x}) + \kappa d^{(y,s,h)}(\mathbf{x})), \tag{A.10}$$

$$0 \leq a^{(y,s,h)}(\mathbf{x}) + \kappa d^{(y,s,h)}(\mathbf{x}), \tag{A.11}$$

³⁴ The endogenous state consists of the asset holdings across age groups. For 150 agents and two assets, this results in a 300-dimensional endogenous state space. Since agents are born without assets, and the beginning-of-period asset holdings of agents aged s^h are the same within a type, irrespective of the health shock, a sufficient endogenous state could be 292-dimensional.

$$(A.12) \quad 0 \leq \mu^{(y,s,h)}(\mathbf{x}),$$

and

$$(A.13) \quad 0 = \sum_{y=1,2} \left(\sum_{s=1}^{s^h-1} d^{(y,s,1)}(\mathbf{x}) + \sum_{s=s^h}^{N-1} (1 - \pi^y) d^{(y,s,1)}(\mathbf{x}) + \pi^y d^{(y,s,2)}(\mathbf{x}) \right),$$

where $r(\mathbf{x}) = f_K(K, L, x_1)$ and $w(\mathbf{x}) = f_L(K, L, x_1)$, with

$$(A.14) \quad K = \sum_{y=1,2} \left(\sum_{s=1}^{s^h-1} k^{(y,s,1)}(\mathbf{x}) + \sum_{s=s^h}^N (1 - \pi^y) k^{(y,s,1)}(\mathbf{x}) + \pi^y k^{(y,s,2)}(\mathbf{x}) \right),$$

$$(A.15) \quad L = \sum_{y=1,2} \left(\sum_{s=1}^{s^h-1} l^{(y,s,1)}(\mathbf{x}) + \sum_{s=s^h}^N (1 - \pi^y) l^{(y,s,1)}(\mathbf{x}) + \pi^y l^{(y,s,2)}(\mathbf{x}) \right).$$

The budget constraint remains unchanged.

As before in our benchmark model, computing an approximation to the FREE in this model means finding an approximation for the $4 \cdot (\tilde{N} - 2) + 1$ equilibrium functions such that the above equations are approximately fulfilled for all exogenous shocks and all values the $2\tilde{N}$ -dimensional endogenous state takes. Besides the larger number of agents and, as a result, the larger number of functions to be approximated and the larger number of equations to be fulfilled, the algorithm remains the same.

Insurance. To study the welfare implication of health insurance in this model, we add additional assets to the model, which pay out contingent on the realization of the health shock. Agents of type y at age $s^h - 1$ can buy an asset, which promises a payout of 1 in case the health shock hits the agent in the following period. The collateral requirements are the same as for the bond. From the perspective of an agent that sells insurance, selling one unit of insurance is indistinguishable from selling π^y units of the bond. Hence, the price $p^y(\mathbf{x})$ of one unit of insurance for type y is pinned down by the bond price and is given by³⁵

$$(A.16) \quad p^y(\mathbf{x}) = \pi^y p(\mathbf{x}).$$

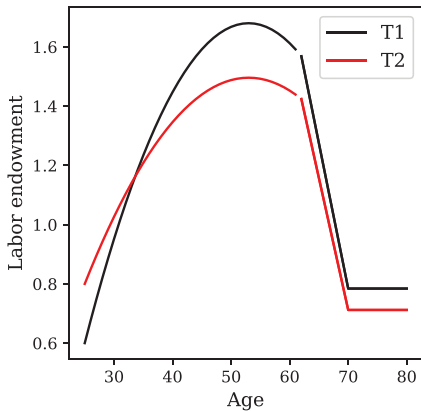
Let $i^y(\mathbf{x})$ denote the units of insurance purchased by age group $s^h - 1$, then the new bond market clearing condition reads

$$(A.17) \quad 0 = \sum_{y=1,2} \left(\pi^y i^y(\mathbf{x}) + \sum_{s=1}^{s^h-1} d^{(y,s,1)}(\mathbf{x}) + \sum_{s=s^h}^{N-1} (1 - \pi^y) d^{(y,s,1)}(\mathbf{x}) + \pi^y d^{(y,s,2)}(\mathbf{x}) \right).$$

The amount of insurance, $i^y(\mathbf{x})$, purchased by an agent of type y and age $s^y - 1$ is pinned down by an additional Euler equation

$$(A.18) \quad p^y(\mathbf{x}) u' \left(c^{(y,s^h-1,1)}(\mathbf{x}) \right) = \beta E_z \left[\pi^y \frac{1}{g^{(y,s^h,2)}} u' \left(\frac{c^{(y,s^h,2)}(\mathbf{x}_+)}{g^{(y,s^h,2)}} \right) \right].$$

³⁵ We can also see the purchase of insurance the following way: the representative agent of type y at age $s^h - 1$ buys π^y units of the bond. The fraction $(1 - \pi^y)$ of households, which do not fall sick the next period, promises the payout of that bond to the fraction π^y of households, which get sick and then receive a payout of $\pi^y + \pi^y \frac{1 - \pi^y}{\pi^y} = 1$.



NOTES: Labor endowment over the life-cycle for type $l = 1$ (in black) and type $l = 2$ (in red) in the models with health shocks. The health-shock occurs at age 62, the year after the line-break.

FIGURE A.6

LABOR ENDOWMENT OVER THE LIFE-CYCLE FOR BOTH TYPES

Here, we do not include any multipliers because we assume that the short-selling constraint on the health insurance never binds. Therefore, we add the relative error in Equation (A.18) directly to the cost function. Note that the cost function for this model has the same form as the cost function of the benchmark model shown in Equation (48), that is, it is given by the mean-squared errors in the relative Euler equations for bond, capital, and insurance as well as the market clearing condition for the bond.

A.4.3 Model parameter setup. In this section, we introduce the parameters, which are new in the OLG model with idiosyncratic shocks. We model $N = 56$ age groups, the health shock occurs at age 62. There are now two, type-specific, profiles of labor endowment over the life cycle (see Figure A.6). The labor profile of type 1 starts lower, rises steeper, and peaks higher. The labor endowment of type $y = 1$ rises by 180% between ages 25 and its peak value at age 53, whereas the labor endowment of type $y = 2$ rises by 87%. The average labor endowment by age group rises by approximately 127%, as in the benchmark model. We chose the probability of the health shock to be $\pi^1 = 0.1$ for type 1 and $\pi^2 = 0.3$ for type 2. The health shock is also more severe for type 2 than for type 1, with $g^{(1,s^h,2)} = 2$ and $g^{(2,s^h,2)} = 3$. The remaining parameters stay the same as in the benchmark model (a summary is given in Table A.1).

A.4.4 Performance of the deep equilibrium net (DEQN). For both cases, the model with and without health insurance, we train the neural network for 100,000 episodes. The chosen hyperparameters are shown in Table A.2. Tables A.3 and A.4, show statistics for the errors in the equilibrium conditions on 10,000 simulated periods after the training was completed. As in the benchmark model, the mean and maximum relative errors in the Euler and KKT conditions remain below 0.1% and 1%, respectively.

A.4.5 Welfare benefits from health insurance. We now make use of our approximated model solution to study the welfare benefits, which arise from the possibility to buy health insurance, that is, an asset with a payoff conditional on receiving the health shock in the context of the model presented in Section A.4.1. To do so, we compare the mean expected utility in the setting with and without insurance. We compare the expected lifetime utility before the agents enter the economy, i.e., before their type has realized, to the expected lifetime utility conditional on the type and conditional on receiving or not receiving the health shock. We convert the difference in lifetime utility to the fraction of consumption an agent would

TABLE A.1
PARAMETERIZATION OF THE MODEL WITH TWO TYPES AND HEALTH SHOCKS (CF., SECTION A.4.1)

	Persistence TFP	Persistence Depreciation	$P(\eta_{t+1} = 1.022 \eta_t = 1.022)$	$P(\delta_{t+1} = 0.08 \delta_t = 0.08)$	$P(\eta_{t+1} = 0.978 \eta_t = 0.978)$	$P(\delta_{t+1} = 0.11 \delta_t = 0.11)$	Borrowing Constraint \underline{a}	Adjustment ζ	Collateral Bond κ	Timing of Shock s^h	Probability of Shock π^1	Probability of Shock π^2	Magnitude of Health Shock $g^{(1,\delta,2)}$	Magnitude of Health Shock $g^{(1,s,2)}$		
Discount Factor β	0.95	2	0.3	{0.978, 1.022}	{0.08, 0.11}	0.905	0.905	0.972	0.700	0.0	0.5	1.1236	38	0.1	0.3	2.3

TABLE A.2
HYPERPARAMETERS CHOSEN TO TRAIN THE DEEP EQUILIBRIUM NET FOR THE MODEL WITH TWO TYPES AND HEALTH SHOCKS

Episodes	Learning Rate α^{learn}	Periods per Episode T	Epochs per Episode N^{epochs}	Mini-Batch Size m	Nodes Hidden Layers	Activations Hidden Layers
1–20,000	1×10^{-5}	10,000	1	64	1,000	relu
20,001–100,000	1×10^{-6}	10,000	1	1,000	1,000	relu
					1,000	relu
					1,000	relu
Episodes	Learning Rate α^{learn}	Periods per Episode T	Epochs per Episode N^{epochs}	Mini-Batch Size m	Nodes Hidden Layers	Activations Hidden Layers
1–18,500	1×10^{-5}	10,000	1	64	1,000	relu
18,501–100,000	1×10^{-6}	10,000	1	1,000	1,000	relu
					1,000	relu
					1,000	relu

NOTE: The top table shows the case without insurance, the bottom table shows the case with insurance.

TABLE A.3
ERRORS IN THE EQUILIBRIUM CONDITIONS FOR THE MODEL WITHOUT HEALTH INSURANCE

	Mean	Max	0.1	10	50	90	99.9
T1, Rel Ee capital [%]	0.025	0.533	0.000	0.002	0.013	0.060	0.310
T1, Rel Ee bond [%]	0.021	0.554	0.000	0.002	0.011	0.050	0.257
T1, KKT capital [$\times 10^{-2}$]	0.001	0.132	0.000	0.000	0.000	0.001	0.039
T1, KKT bond [$\times 10^{-2}$]	0.003	0.150	0.000	0.000	0.000	0.011	0.052
T2, Rel Ee capital [%]	0.028	0.844	0.000	0.002	0.014	0.068	0.345
T2, Rel Ee bond [%]	0.024	0.881	0.000	0.002	0.012	0.057	0.321
T2, KKT capital [$\times 10^{-2}$]	0.000	0.140	0.000	0.000	0.000	0.001	0.013
T2, KKT bond [$\times 10^{-2}$]	0.004	0.196	0.000	0.000	0.000	0.013	0.072
Market clearing [$\times 10^{-2}$]	0.193	1.679	0.001	0.056	0.169	0.362	1.003
Market clearing (normalized) [$\times 10^{-2}$]	0.000	0.004	0.000	0.000	0.000	0.001	0.002

NOTE: The first four rows show the equilibrium conditions for type 1 agents, rows 5–8 show the conditions for type 2 agents. The first two rows show the relative Euler equation errors (Rel Ee in %) on 10,000 simulated periods. The next two rows show the errors in the KKT conditions ($\times 10^{-2}$). Rows 9 and 10 show the errors in the market clearing conditions ($\times 10^{-2}$). In row 10, the error in the market clearing condition for the bond is divided by aggregate production. The columns show the mean and the max errors as well as the 0.1st, 10th, 50th, 90th, and 99.9th percentile.

TABLE A.4
ERRORS IN THE EQUILIBRIUM CONDITIONS FOR THE MODEL WITH HEALTH INSURANCE

	Mean	Max	0.1	10	50	90	99.9
Rel Ee capital [%]	0.024	0.528	0.000	0.002	0.013	0.060	0.324
Rel Ee bond [%]	0.021	0.608	0.000	0.002	0.012	0.051	0.255
KKT capital [$\times 10^{-2}$]	0.000	0.127	0.000	0.000	0.000	0.000	0.000
KKT bond [$\times 10^{-2}$]	0.005	0.104	0.000	0.000	0.000	0.000	0.021
Market clearing [$\times 10^{-2}$]	0.053	0.333	0.000	0.017	0.047	0.089	0.252
Market clearing (normalized) [$\times 10^{-2}$]	0.000	0.002	0.000	0.000	0.000	0.000	0.001

NOTE: The first five rows show the equilibrium conditions for type 1 agents, rows 6–9 show the conditions for type 2 agents. The first three rows show the relative Euler equation errors (Rel Ee in %) on 10,000 simulated periods. The next two rows show the errors in the KKT conditions ($\times 10^{-2}$). Rows 11 and 12 show the errors in the market clearing conditions ($\times 10^{-2}$). In row 12, the error in the market clearing condition for the bond is divided by aggregate production. The columns show the mean and the max errors as well as the 0.1st, 10th, 50th, 90th, and 99.9th percentile.

TABLE A.5
MEAN LIFETIME UTILITY IN THE MODEL WHERE NO INSURANCE IS AVAILABLE

	Cond. on Type						
	Cond. on Health Shock						
	Unconditional	T1	T2	T1, sick	T1, healthy	T2, sick	T2, healthy
Mean lifetime utility	-18.206	-18.533	-17.878	-18.609	-18.524	-17.992	-17.830
Consumption factor [%] (compared to unconditional)	0.0	-1.8	+1.8	-2.2	-1.7	+1.2	+2.1

NOTE: Mean lifetime utility in the model where no insurance is available, computed from 100,000 simulations of 150 time periods. The first row shows the mean lifetime utility; the second row shows the percentage of extra consumption the mean agent would need to receive in order to attain the same mean lifetime utility as lucky/unlucky agents, who received specific shocks.

additionally need to receive in every state of the world in order to attain the same lifetime utility in both cases. This factor, denoted by q , can be computed noting that for CRRA utility with coefficient of relative risk aversion γ , the expected lifetime utility scales by a factor $(1+q)^{1-\gamma}$ if consumption in every state is scaled by a factor $(1+q)$: $V((1+q)c) = (1+q)^{1-\gamma}V(c)$.

Table A.5 shows the lifetime utility in the economy without insurance. As shown in Table A.5, type 2 agents who do not receive the health shock attain the highest mean

TABLE A.6
MEAN LIFETIME UTILITY IN THE MODEL WHERE INSURANCE IS AVAILABLE

	Conditional on Type						
	Conditional on Health Shock						
	Unconditional	T1	T2	T1, sick	T1, healthy	T2, sick	T2, healthy
Mean lifetime utility	-18.204	-18.531	-17.877	-18.569	-18.527	-17.933	-17.852
Consumption factor [%] (compared to unconditional)	0.0	-1.8	+1.8	-2.0	-1.7	+1.5	+2.0

NOTE: Mean lifetime utility in the model where insurance is available, computed from 100,000 simulations of 150 time periods. The first row shows the mean lifetime utility. The second row shows the percentage of extra consumption the mean agent would need to receive in order to attain the same mean lifetime utility as lucky/unlucky agents, who received specific shocks.

lifetime utility. The difference to the unconditional average corresponds to approximately 2.1% of consumption. Furthermore, the table shows that the type of shock at birth has a larger influence on the mean attained lifetime utility than the health shock. The difference in mean received lifetime utility between agents born as type 1 and agents born as type 2 corresponds to 3.6% of consumption. The difference between receiving and not receiving the shock is approximately 0.5% and 0.9% for type 1 and 2, respectively.

Table A.6 shows the same numbers for the economy in which health insurance is available. The effect on mean attained lifetime utility is not significant when looking at the unconditional average, or the average conditional on the type. This may be due to the fact that most of the lifetime utility is accumulated in earlier years, before the health shock hits. Since the health shock occurs in the 38th modeled period (at age 62), utility attained in the remaining periods is discounted with $\beta^{37} \approx 0.15$ and more when computing attained lifetime utility.

The difference in attained lifetime utility between agents who receive the health shock and those who do not is halved through the availability of insurance. Focusing on the utility accumulated from age 62 (the age at which the health shock occurs) and onward, the difference in utility attained by agents who fell sick and who stayed healthy corresponds to -6.6% of consumption for type 1 agents and to -11.2% of consumption for type 2 agents in the model without health insurance. In the model with health insurance, the differences reduce to -3.4% and -5.8%. In our stylized model, the ability to buy health insurance hence has the most significant impact on older agents.

A.5 Application with a Continuum of Agents, and Aggregate and Idiosyncratic Shocks. Our method is generic by construction and can be applied to a wide range discrete-time dynamic (stochastic) models. To illustrate this, we apply the DEQN framework in a setting with three additional innovations. That is, we solve a model with (i) a continuum of ex ante identical agents, (ii) per-period aggregate and idiosyncratic shocks, and (iii) where the agents endowed with recursive (Epstein–Zin) preferences. From a computational perspective, Epstein–Zin preferences require us to approximate the value function in addition to the policy function. We show how our method is able to handle this additional requirement, simply by adding the Bellman equation to the loss function. Conveniently, most parts of the algorithm (and the code) hence remain unchanged when modeling recursive preferences. We model six discrete aggregate shocks and two regimes of aggregate uncertainty. From an economic perspective, Epstein–Zin preferences are of particular interest in this setting since they allow for a separation between the risk aversion and the intertemporal elasticity of substitution.

Following the approach of Young (2010), we encode the distribution of asset holdings across households for each idiosyncratic shock into a histogram and use nonstochastic simulation to obtain its transition given the households' policies and a sequence of exogenous

aggregate shocks.³⁶ This approach has the advantage that it allows the precise evaluation of the expectation operators³⁷ and it gives the network more direct access to the wealth distribution and hence to economics that it would otherwise have to learn. For example, if the aggregate state is encoded as an array of agents' asset holdings and simulated shocks, the ordering of the agents should not affect the aggregate state. Encoding the distribution as a histogram avoids this issue, facilitating the learning of the policy.³⁸

A.5.1 The Bewley (1977) model with aggregate shocks. Similar to the benchmark model, one of A possible aggregate shocks realizes every period. We denote the aggregate shock realizing in period t by a_t and a history of aggregate shocks up to period t by $a^t = (a_0, \dots, a_t)$. The aggregate labor income in the economy, denoted by $W_t = W(a_t)$, is exogenous and only depends on the aggregate shock. In contrast to the previous model, we now consider a continuum of agents facing idiosyncratic shocks. Let there be I idiosyncratic shocks, where i_t denotes a particular shock in period t , and i^t denotes a history of shocks. We assume that both shocks follow a first-order Markov process with finite support \mathcal{A} and \mathcal{I} , and denote the associated transition matrices with Π^A and Π^I , respectively.

The share of aggregate labor endowment owned by an agent, which depends only on the realization of the idiosyncratic shock, is denoted by $\eta_t = \eta(i_t)$. Let $w_t = w(a_t, i_t)$ denote the exogenous labor income of an agent in period t with aggregate shock a_t and idiosyncratic shock i_t , where $w_t = W_t \eta_t$. Let b_t denote an agent's beginning-of-period bond holdings. We assume the bond is in unit net supply. Furthermore, let \underline{b} denote the exogenous borrowing constraint such that $b_t \geq \underline{b}$. The price of the bond in period t is denoted by p_t . Aggregate consumption is given by $C_t = W_t + 1 - p_t$.

To solve this model recursively, a state of the economy has to be defined on which the policy, value, and price functions depend. First, let μ_t denote the distribution, such that $\mu(i_t, b_t)$ denotes the mass of agents with idiosyncratic shock i_t and asset holding b_t . Then, the aggregate state of the economy is given by the exogenous shock a_t , and the distribution of agents across asset holdings and idiosyncratic shocks, $x_t^{\text{agg}} = (a_t, \mu_t)$. The idiosyncratic state of an agent is given by $x_t^{\text{id}} = (i_t, b_t)$. Aggregate quantities, that is, the bond price $p_t = p(x_t^{\text{agg}})$, are then functions of the aggregate state alone. The agents' policy and value functions are functions of the aggregate and idiosyncratic state $x_t := (x_t^{\text{id}}, x_t^{\text{agg}})$, that is, $b_{t+1} = b(x_t)$ and $V_t = V(x_t)$. Then, the agent's value function is given by

$$(A.19) \quad V(x_t) = \max_{b_{t+1}} \left((1 - \beta)c_t^{1-\rho} + \beta \chi_t(V(x_{t+1}))^{1-\rho} \right)^{\frac{1}{1-\rho}}, \text{ where}$$

$$(A.20) \quad \chi_t(V_{t+1}) := E_t[V_{t+1}^{1-\sigma}]^{\frac{1}{1-\sigma}}, \text{ and}$$

$$(A.21) \quad c_t = w_t + b_t - b_{t+1}p_t.$$

The corresponding KKT conditions are given by

$$(A.22) \quad p_t c_t^{-\rho} = \beta E_t \left[c_{t+1}^{-\rho} \left(\frac{V_{t+1}}{\chi_t(V_{t+1})} \right)^{\rho-\sigma} \right] + \lambda_t,$$

$$(A.23) \quad b_{t+1} \geq \underline{b},$$

³⁶ Alternatively, one could solve models with idiosyncratic and aggregate risk by taking a large but finite number of agents (Maliar et al., 2021). The aggregate state is then encoded as an array of individual asset holdings and shocks, and the transition is obtained by stochastic simulation.

³⁷ Note that an approximation is still involved when sorting asset holdings into a histogram.

³⁸ An alternative way to tackle this issue would be to exploit the symmetry explicitly, for example, through the design of the state, the sampling method, or the architecture of the neural network. This approach is laid out in recent work by Kahou et al. (2021) for models with (finitely) many agents.

$$(A.24) \quad \lambda_t \geq 0,$$

$$(A.25) \quad \lambda_t(b_{t+1} - \underline{b}) = 0.$$

To approximate these functions numerically, we follow Reiter (2009) and Young (2010) and encode the distribution with a histogram of finitely many discrete values. We define the histogram on a grid of N_b equidistant points $(b_1^{\text{grid}}, \dots, b_{N_b}^{\text{grid}})$. We set the lower bound of the histogram to $b_0^{\text{grid}} = \underline{b}$ and choose $b_{N_b}^{\text{grid}}$ large enough, such that the mass in that histogram bin remains small when simulating the evolution of the histogram. Then, we denote the histogram-based approximation of the distribution by $\mu_t \in \mathbb{R}^{I \cdot N_b}$. The dimension of the approximated aggregate state and the state governing the agents' decisions are then given by $N_{\text{agg}} = I \cdot N_b + 1$ and $N_x = N_{\text{agg}} + 2$, respectively.

Since one usually needs more than 20 histogram bins to approximate the wealth distribution accurately, the state space becomes high-dimensional. Many standard methods to compute equilibria globally are not able to handle such a high-dimensional state space and hence have to resort to techniques which find a low-dimensional statistic for the distribution, which is sufficient to predict prices accurately (along the lines of the seminal work by Krusell and Smith, 1998). For neural networks on the other hand, inputs with up to thousands of variable do not pose a challenge. Furthermore, a deep neural network can learn to extract the relevant features of the distribution, which it then uses in the later layers for prediction, by itself. Consequently, our method naturally allows us to pass the full discretized distribution as input to the neural network.

FREE.

DEFINITION A.2 (FUNCTIONAL RATIONAL EXPECTATIONS EQUILIBRIUM). A FREE consists of equilibrium functions $\{V(\cdot), b(\cdot), \lambda(\cdot), p(\cdot)\}$, fulfilling conditions (A.26)–(A.34). $V : \mathcal{A} \times \mathcal{I} \times \mathbb{R}^{N_x-2} \rightarrow \mathbb{R}$ denotes the value function, $b : \mathcal{A} \times \mathcal{I} \times \mathbb{R}^{N_x-2} \rightarrow \mathbb{R}$ denotes the investment function, $\lambda : \mathcal{A} \times \mathcal{I} \times \mathbb{R}^{N_x-2} \rightarrow \mathbb{R}$ denotes the KKT multiplier function, and $p : \mathcal{A} \times \mathbb{R}^{N_{\text{agg}}-1} \rightarrow \mathbb{R}$ denotes the price function, such that for all states $\mathbf{x} = (\mathbf{x}^{\text{id}}, \mathbf{x}^{\text{agg}}) = (i, b, a, \mu)$, we have that:

$$(A.26) \quad V(\mathbf{x}) = ((1 - \beta)c^{1-\rho} + \beta\chi(V(\mathbf{x}_+))^{1-\rho})^{\frac{1}{1-\rho}},$$

$$(A.27) \quad p(\mathbf{x}^{\text{agg}})c^{-\rho} = \beta E \left[c_+^{-\rho} \left(\frac{V(\mathbf{x}_+)}{\chi(V(\mathbf{x}_+))} \right)^{\rho-\sigma} \right] + \lambda(\mathbf{x}),$$

$$(A.28) \quad c = w(a, i) + b - p(\mathbf{x}^{\text{agg}})b(\mathbf{x}),$$

$$(A.29) \quad \mathbf{x}_+ = [i_+, b(\mathbf{x}), a_+, \mu_+]^T,$$

$$(A.30) \quad \mu_+ \text{ consistent with } \mathbf{x}^{\text{agg}} \text{ and } b(\cdot),$$

$$(A.31) \quad 1 = \sum_{i=1}^I \sum_{j=1}^{N_b} \mu_{(i-1)N_b+j} b([i, b_j^{\text{grid}}, \mathbf{x}^{\text{agg}}]^T),$$

$$(A.32) \quad 0 = \lambda(\mathbf{x})(b(\mathbf{x}) - \underline{b}),$$

$$(A.33) \quad 0 \geq \lambda(\mathbf{x}),$$

$$(A.34) \quad 0 \geq (b(\mathbf{x}) - \underline{b}).$$

Equations (A.26) and (A.27) ensure that the Bellman and Euler equation are satisfied. Equation (A.28) gives the budget constraint. Equations (A.29) and (A.30) give the transition of that aggregate state, implied by policy $b(\cdot)$. Equation (A.31) ensures that the market clears. Finally, Equations (A.32)–(A.34) ensure that the remaining KKT conditions are satisfied.

Approximation with DEQNs. We approximate the aggregate and idiosyncratic equilibrium functions using two neural networks.³⁹ The neural network that approximates the bond price is denoted with $\mathcal{N}_{\rho_p}^p$, where ρ_p denotes its trainable parameters. The neural network that approximates the value functions, policies, and multipliers, is denoted by $\mathcal{N}_{\rho_{id}}^{id}$ with trainable parameters ρ_{id} . We define \hat{V} , \hat{b} , $\hat{\lambda}$, and \hat{p} as the corresponding approximations resulting from the neural networks:

$$(A.35) \quad \mathcal{N}_{\rho_p}^p(\mathbf{x}^{agg}) := \hat{p}(\mathbf{x}^{agg}) \approx p(\mathbf{x}^{agg}),$$

$$(A.36) \quad \mathcal{N}_{\rho_{id}}^{id}(\mathbf{x}) := [\hat{V}(\mathbf{x}), \hat{b}(\mathbf{x}), \hat{\lambda}(\mathbf{x})]^T \approx [V(\mathbf{x}), b(\mathbf{x}), \lambda(\mathbf{x})]^T.$$

Loss function. As before, we design the neural network such that Equations (A.33) and (A.34) are always satisfied.⁴⁰ Furthermore, we ensure the budget constraint holds by computing consumption directly using Equation (A.28). Similarly, we obtain μ_+ from the current policy, ensuring conditions (A.29) and (A.30) always hold precisely. We define the implicitly approximated variables as $\hat{c}(\mathbf{x})$, $\hat{\mu}_+(\mathbf{x})$, and $\hat{\mathbf{x}}_+$. The remaining equilibrium conditions (i.e., (A.26), (A.27), (A.31), and (A.32)) are encoded in a loss function which is minimized by the neural network. For a given state \mathbf{x} , the errors in these conditions are given by

$$(A.37) \quad \text{err}_{\mathbf{x}}^{\text{Bellman}}(\rho_p, \rho_{id}) := \frac{\left((1 - \beta) \hat{c}(\mathbf{x})^{1-\rho} + \beta \chi (\hat{V}(\hat{\mathbf{x}}_+))^{1-\rho} \right)^{\frac{1}{1-\rho}}}{\hat{V}(\mathbf{x})} - 1,$$

$$(A.38) \quad \text{err}_{\mathbf{x}}^{\text{Euler}}(\rho_p, \rho_{id}) := \frac{\left(\frac{\mathbb{E} \left[\hat{c}(\hat{\mathbf{x}}_+)^{-\rho} \left(\frac{\hat{V}(\hat{\mathbf{x}}_+)}{\chi(\hat{V}(\hat{\mathbf{x}}_+))} \right)^{\rho-\sigma} \right] + \hat{\lambda}(\mathbf{x})}{\hat{p}(\mathbf{x}^{agg})} \right)^{-\frac{1}{\rho}}}{\hat{c}(\mathbf{x})} - 1,$$

$$(A.39) \quad \text{err}_{\mathbf{x}^{agg}}^{\text{mc}}(\rho_p, \rho_{id}) := \sum_{i=1}^I \sum_{j=1}^{N_b} \mu_{(i-1)N_b+j} \hat{b}([i, b_j^{grid}, \mathbf{x}^{agg}]^T) - 1,$$

$$(A.40) \quad \text{err}_{\mathbf{x}}^{\text{slack}}(\rho_p, \rho_{id}) := (\hat{b}(\mathbf{x}) - \underline{b}) \hat{\lambda}(\mathbf{x}).$$

For a given training set of states, $\mathcal{D}_{\text{train}}$, we define the loss function as the mean-squared error of the equilibrium conditions:

$$(A.41) \quad \ell_{\mathcal{D}_{\text{train}}}(\rho_p, \rho_{id}) := \frac{1}{|\mathcal{D}_{\text{train}}|} \sum_{\mathbf{x} \in \mathcal{D}_{\text{train}}} (\text{err}_{\mathbf{x}}^{\text{Bellman}}(\rho_p, \rho_{id})^2 + \text{err}_{\mathbf{x}}^{\text{Euler}}(\rho_p, \rho_{id})^2 + \text{err}_{\mathbf{x}}^{\text{mc}}(\rho_p, \rho_{id})^2 + \text{err}_{\mathbf{x}}^{\text{slack}}(\rho_p, \rho_{id})^2).$$

Simulating states. Given the histogram starting point μ_0 and a sequence of aggregate shocks (a_0, \dots, a_{T-1}) , we use the policy encoded by the policy network $\mathcal{N}_{\rho_{id}}^{id}$ to obtain a sequence of histograms $(\mu_0, \dots, \mu_{T-1})$. We combine the sequence of shocks and histograms to

³⁹ With aggregate equilibrium functions we refer to functions, which predict aggregate quantities and only depend on the aggregate state of the economy. In this model, the only aggregate equilibrium function is the function for the bond price. With idiosyncratic equilibrium functions, we refer to functions which depend, on the aggregate as well as the idiosyncratic state. In this model, the idiosyncratic equilibrium functions are the households' policy-, multiplier- and value functions.

⁴⁰ That is, we use a softplus activation function in the network's output layer and approximate $b - \underline{b}$ instead of b .

obtain a sequence of aggregate states $(\mathbf{x}_0^{\text{agg}}, \dots, \mathbf{x}_{T-1}^{\text{agg}})$. Then, we construct a training set $\mathcal{D}_{\text{train}}$ by randomly drawing N idiosyncratic states for each of the T aggregate states.⁴¹

Training with two neural networks. The DEQN is trained analogously to the previous model. That is, our algorithm iterates between a simulation and a network training phase, updating the neural networks' parameters using a variant of mini-batch stochastic gradient descent. Using two separate neural networks does not pose an additional challenge since a single forward and backward pass allows us to obtain the gradients of the loss function with respect to the parameters ρ_p and ρ_{id} , simultaneously. We then use the gradients to update the parameters as previously described.

A.5.2 Parameters and performance of the DEQN.

Model parameters. We chose to model six discrete aggregate shocks and two discrete idiosyncratic shocks. We model two aggregate uncertainty regimes; three discrete shocks for each of the regimes. This is an interesting application for our method since the literature on uncertainty shocks often requires global and nonlinear solution methods (see, e.g., Bloom et al., 2018; Fernández-Villaverde and Guerron-Quintana, 2020). Furthermore, it is a setting where Epstein-Zin utility is of special interest since it allows for a distinction between the risk-aversion and the elasticity of intertemporal substitution. The aggregate labor income for the six aggregate shocks by $W = [0.95, 1.00, 1.05, 0.95, 1.00, 1.05]^T$, where the first three shocks correspond to times of lower aggregate uncertainty and the second three shocks correspond to times of higher aggregate uncertainty. The transition matrix to transition between the aggregate labor income $W = [0.95, 1.00, 1.05]^T$ is given by $\Pi^{A,l}$ in times of lower and by $\Pi^{A,h}$ in times of higher aggregate uncertainty. The probability to switch uncertainty regimes is denoted by $\Pi^{A,\text{switch}}$. The parameter values are given by

$$\Pi^{A,l} = \begin{bmatrix} 0.70 & 0.20 & 0.10 \\ 0.15 & 0.70 & 0.15 \\ 0.10 & 0.20 & 0.70 \end{bmatrix}, \quad \Pi^{A,h} = \begin{bmatrix} 0.75 & 0.10 & 0.15 \\ 0.20 & 0.60 & 0.20 \\ 0.15 & 0.10 & 0.75 \end{bmatrix}, \quad \Pi^{A,\text{switch}} = \begin{bmatrix} 0.8 & 0.2 \\ 0.2 & 0.8 \end{bmatrix}$$

$$(A.42) \Rightarrow \Pi^A = \begin{bmatrix} 0.8\Pi^{A,l} & 0.2\Pi^{A,l} \\ 0.2\Pi^{A,h} & 0.8\Pi^{A,h} \end{bmatrix} \in \mathbb{R}^{6 \times 6}.$$

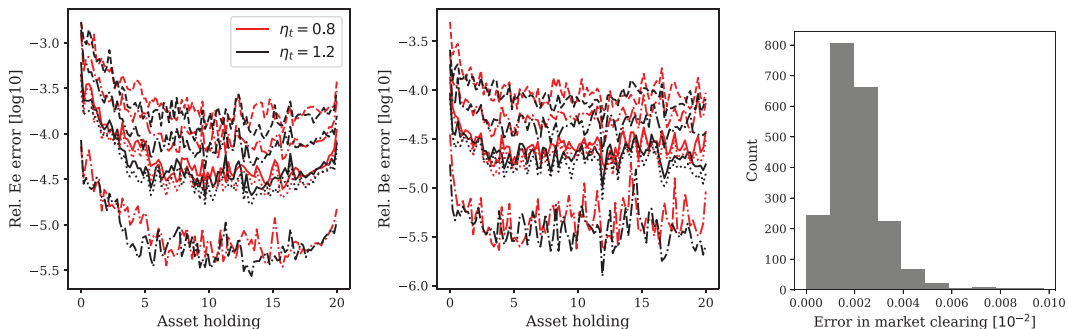
As in Fang (2021), the difference between the transition probabilities is that in times with higher aggregate uncertainty, the probability to transition into the two extreme realizations $W = 0.95$ and $W = 1.05$ is larger (by 5%, respectively). On the other hand, the probability of transitioning into the middle realization $W = 1.00$ is smaller (by 10%, respectively). For the idiosyncratic shocks we chose $\eta = [0.8, 1.2]^T$ with transition probabilities

$$(A.43) \quad \Pi^I = \begin{bmatrix} 0.7 & 0.3 \\ 0.3 & 0.7 \end{bmatrix}.$$

For the preferences, we chose a constant relative risk aversion of $\sigma = 8$ and an intertemporal elasticity of substitution of $\frac{1}{\rho} = 0.5$. The patience is chosen to be $\beta = 0.95$ and the borrowing constraint is at $b = 0$.

Neural networks. To pass the distribution over asset holdings to the neural networks, we discretize it to 100 points for each of the two idiosyncratic shocks. The discretized distribution is hence encoded as a 200-dimensional vector. The aggregate state further consists of the

⁴¹ We draw the idiosyncratic shock randomly according to its unconditional probability, and the idiosyncratic asset holding uniformly random from the interval $[b_1^{\text{grid}}, b_{N_b}^{\text{grid}}]$.



NOTES: The red lines show statistics for the bad idiosyncratic shock and black lines statistics for the good idiosyncratic shocks. The statistics are based on 2,048 simulated aggregate states, for each of the asset holdings and each of the idiosyncratic shocks. The dash-dotted lines show to 10th and 90th percentiles, the dotted lines the median, the solid lines the mean, and the dashed line the 99.9th percentile.

FIGURE A.7

RELATIVE ERRORS IN THE EULER (LEFT PANEL) AND BELLMAN (MIDDLE PANEL) EQUATIONS

aggregate shock and two additional (redundant) indicators, one for the value of the wage income and one for the uncertainty regime, rendering the aggregate state 203-dimensional. The idiosyncratic state additionally includes an indicator for the idiosyncratic shock and the asset holding. Hence, the input layer consists of 203 nodes for the price network and 205 nodes for the policy network. Each of the networks further consists of two relu activated hidden layers with 500 nodes each. The price network has a single softplus activated node in the output layer, which predicts the price. The policy network has three softplus activated nodes in the output layer predicting the bond purchase, the value, and the multiplier associated with the borrowing constraint.

Simulation. In each episode, we simulate 2,048 aggregate states (16 tracks, 128 periods each). For each simulated aggregate state and each of the two discrete idiosyncratic shocks, we randomly draw 20 values for the idiosyncratic asset holding uniformly between the lower and a sufficiently high upper bound to construct the idiosyncratic states.⁴² This results in a total of 81,920 idiosyncratic and 2,048 aggregate states per episode.

Training. We train the neural network for 30,000 episodes with a learning rate of 1×10^{-5} and a batch size of 126. Subsequently, we train it for an additional 2,000 episodes with a batch size of 1,024 and a learning rate of 1×10^{-6} .

Performance. Figure A.7 shows the accuracy after training the neural networks for 32,000 episodes. The mean relative errors in the Euler and Bellman equation are below 10^{-3} , and even the 99.9th percentile is below 10^{-2} for the Euler equation and below 10^{-3} for the Bellman equation. The error in the market clearing condition for the bond is of the order 10^{-4} . The errors in the KKT conditions are low as well.

A.6 Background on Neural Networks

A.6.1 Why neural networks. In light of contemporary developments, neural networks have become a popular choice of method for a wide range of applications. The availability of large data sets, improvements in hardware, and algorithmic advancements have made neural networks very competitive for data processing tasks. For computational tasks, neural networks

⁴² The lower bound on the asset holding is 0 and the upper bound is 20.

TABLE A.7
COMPARISON OF FEATURES OF DIFFERENT APPROXIMATION METHODS

Approximation Method	High-Dimensional Input	Can Resolve Local Features Accurately	Irregularly Shaped Domain	Large Amount of Data
Polynomials	✓	✗	✓	✓
Splines	✗	✓	✗	✓
Adaptive (sparse) grids	✓	✓	✗	✓
Gaussian processes	✓	✓	✓	✗
Deep neural networks	✓	✓	✓	✓

possess features, which, along with their theoretical properties (see Appendix A.7.1), offer a decisive advantage. In Table A.7, we summarize these features in relation to other standard approximation methods. The correct choice of approximator depends on the features needed to address a given problem. For problems requiring all of the listed features, neural networks, specifically DEQNs, offer a promising approach.

A densely connected feed forward neural network is a sequence of matrix–vector multiplications, vector–vector additions, and elementwise applications of the activation functions. Hence, the computational complexity of an evaluation of the neural network is inherited from these operations. Furthermore, the computational complexity of computing the gradients with automatic differentiation is of similar magnitude to that of a single evaluation of the neural network. Therefore, in both training and inference, neural networks can handle very high-dimensional inputs without becoming prohibitively slow.

In addition, neural networks can handle large amounts of input data and irregularly shaped domains and can approximate functions with kinks and ridges.⁴³ In contrast, polynomials are well suited for approximating high-dimensional smooth functions on irregularly shaped domains but have difficulties resolving kinked features; adaptive (sparse) grids can approximate functions with local features using a large amount of high-dimensional data but are inefficient on irregular domains, and Gaussian processes can approximate high-dimensional functions with local features on irregularly shaped domains, but become prohibitively slow when using many training points. Given the presence of all of the previous features, as is the case in the models presented above, neural networks offer a suitable approximator.

We do not believe that deep learning is a silver bullet that should always be used. Ultimately, the appropriate solution method depends on the economic model at hand, and its determination requires expert knowledge of said model. For example, if the dimensionality of the state space is moderate (e.g., 10 dimensions or fewer) and if the problem can trivially be mapped to a hypercubic domain, grid-based methods, such as adaptive sparse grids (see, e.g., Brumm et al., 2021, and references therein), may be the preferred choice.

A.6.2 Neural networks and the curse of dimensionality. There is a broad body of literature experimentally demonstrating that neural networks ameliorate the curse of dimensionality in a variety of settings (see, e.g., Bengio and Bengio, 2000; Han et al., 2018; Becker et al., 2019; Beck et al., 2021). Furthermore, formal proofs are available for specific settings (see, e.g., Bach, 2017; Grohs et al., 2018; Hutzenthaler et al., 2020). Intuitively, we think the advantages of deep learning stem from a combination of local adaptivity, the ability to optimally

⁴³ To approximate local features, such as kinks, accurately, it is important that the approximating function can fit strong nonlinearities in some areas of the domain without deteriorating the approximation quality in others. While some grid-based methods, like adaptive sparse grids or linear interpolation from a dense grid, can resolve local features well, it is not the case for polynomial interpolation (approximating a kinked function by interpolating points with global polynomials often leads to undesired “waves”—known as Runge’s phenomenon—which also occur away from the kink).

learn lower-dimensional representations of the input data, as well as its ability to exploit recent advances in computing hardware (see, e.g., Goodfellow et al., 2016, for a general introduction to deep learning).

A.6.3 Mini-batches, stochastic gradient descent, and the learning rate. Gradient descent is a first-order optimization method in which a loss function is minimized by updating an algorithm's parameters in the decreasing direction of the loss function's gradient. There are many variations and extensions to the gradient descent optimization algorithm that help accelerate and stabilize training. We briefly outline those used in this article: namely, the learning rate, mini-batch gradient descent, and the Adam optimizer.

The learning rate, which corresponds to the step size taken relative to the gradient (see Equation (49)), is one hyperparameter that can be tuned to facilitate training. We found it helpful to use a larger learning rate at the beginning of training and decrease it toward the end. A large learning rate allows for the quick traversal of the loss function while the parameters are “far” from the optimum. On the other hand, a small learning rate enables a tighter convergence around the optimum. By setting the learning rate too high, we can overshoot and oscillate around the optimum or land in a region where the parameters generate economically nonsensical values (see Appendix A.7.3). Contrarily, by setting the learning rate too low, training can be unnecessarily slowed or halted entirely if nonlinear features of the loss function cannot be overcome.

Mini-batch stochastic gradient descent is another method that can be used. Instead of computing the gradients on the full data set, which can be computationally expensive, the gradients are estimated on a subset of randomly sampled data (without replacement), that is, on a *mini-batch*. With mini-batches, the estimates of the gradients are computationally cheaper than with gradient descent. Furthermore, we can perform multiple updates of the network parameters in a single epoch. Finally, the variance of the estimates can be controlled by fine-tuning the mini-batch size. That is, the smaller the mini-batches, the higher the variance of the estimates.

In addition, various gradient descent optimizers stabilize the training of neural networks by continually adapting the learning rate and gradient.⁴⁴ To this end, the Adam optimizer (Kingma and Ba, 2014) uses decayed momentum and a decayed adaptive learning rate. That is, *momentum* updates the parameters using a weighted moving average of the previous and current gradients, thereby favoring steps that occur in a consecutively consistent direction. In Adam, the learning rates are adapted through the normalization by a weighted average of the gradient's second moment.

A.6.4 Supervised versus unsupervised learning. Supervised learning is generally based on a data set of label data. A labeled data set is a set of input–output pairs $\mathcal{D} = \{(x_i, y_i)\}_{i=1}^N$, where y_i is the desired output for input x_i . One then trains the neural network so that the neural network's predictions for inputs x_i are close to the corresponding label y_i . In our algorithm, the inputs are states of the economy, and the outputs are the endogenous variables. Thus, we do not have labels y_i available. For example, in the benchmark model, the inputs x_i are given by the exogenous shock and the capital and bond holdings by age group. The desired output would be the equilibrium savings in capital, the savings in bond, the associated KKT multipliers, as well as the bond price. The labels y_i would be the true equilibrium values for those variables. Hence, we do not have access to labeled data. Since we train the neural network without the need for labeled data, our algorithm is unsupervised. Generally, the availability of label data is one of the main bottlenecks for training neural networks to high accuracy. Our algorithm circumvents this bottleneck because the loss function we construct does not need labels to be evaluated. This is crucial because it enables us to generate a large amount

⁴⁴ See, for example, Ruder (2016) for an extensive overview of neural network optimization methods.

of training data cheaply.⁴⁵ For more information on supervised and unsupervised learning, see James et al. (2013).

A.7 Practical Issues.

A.7.1 Practical issues of DEQNs. In this section, we briefly describe some of the challenges our solution method faces in practical applications. A significant drawback of deep neural networks is, in general, that both the theoretical convergence rates as well as why and when deep neural networks can ameliorate or break the curse of dimensionality are poorly understood. Even though substantial progress has recently been made (see, e.g., Montanelli and Du, 2019), methods such as ours still heavily depend on numerical experimentation. Such shortcomings are consequently reflected in our solution framework.

A key issue common to the field of deep learning is that the choice of hyperparameters is crucial for the success of a learning algorithm. Nevertheless, the choice is often ad hoc or based on prior research experience. The development of frameworks to automate the process of hyperparameter optimization is underway and may help mitigate this issue in future research.

Our proposed algorithm is not guaranteed to converge to a satisfactory solution. Although we often obtain very good results, we also encountered instances for which the algorithm seems to converge despite significant errors in the equilibrium conditions. In this regard, the iteration between stochastic simulation and variants of mini-batch gradient descent on simulated data may pose a particular challenge: if the simulated states happen to be too far from states the neural network was previously trained on, the approximation on the simulated states may be poor and thus lead to very large Euler equation errors or punishment factors due to, for example, policies implying negative consumption. When evaluated on such states, the gradients of the loss function with respect to the neural network's parameters may be so large that all the previous learning progress is overwritten. This may lead to vastly different policies, which in turn, may amplify the problem as consecutively simulated states may be far from the states the neural network was trained on previously. In some instances, this problem can be mitigated by choosing a lower learning rate or larger batch sizes. Alternatively, in some cases, it may be possible to determine an approximation of the ergodic set of states without simulation, which can be sampled accordingly. However, as previously stated, we consider the approximation on simulated states an essential feature of our algorithm and, hence, it would be desirable to find sufficient conditions for convergence.

Furthermore, in some cases, we found that several KKT multipliers were “stuck” very close to zero for many episodes of training despite a positive value yielding much smaller errors in the equilibrium conditions. Recall that we use a softplus activation function in the output layer to ensure that the nonnegativity constraints on the KKT multipliers are always satisfied.⁴⁶ A downside of using a softplus activation function is that its derivative decreases exponentially as its output approaches zero. Consequently, it can take a long time for small gradients to accumulate and lead to a larger output. If this problem occurs, a possible solution may be to use other (less standard) nonnegative activation functions in the output layer,⁴⁷ or to use a different strategy to encode the KKT conditions into the loss function.⁴⁸

From a more theoretical viewpoint, deep learning essentially relies on two pillars: the universal approximator property of the neural networks (see, e.g., Hornik et al., 1989) and

⁴⁵ In the case of standard value function iteration, one would use the solution to the maximization problem to update the approximation of the value function. That solution could be interpreted as temporary labeled data to “train” (update) the value function. The generation of these temporary labels involves solving a maximization problem and is hence computationally costly.

⁴⁶ The softplus activation function is given by $\sigma(x) = \ln(1 + e^x)$.

⁴⁷ For example, $\sigma(x) = x^2$.

⁴⁸ Kotlikoff et al. (2021a) and Maliar et al. (2021), for example, make use of the Fischer–Burmeister function to replace the KKT conditions.

stochastic gradient descent, which is a robust estimate of the classical gradient descent method and thus is only guaranteed to find the global optimal parameters of the model for convex problems (see, e.g., Deisenroth et al., 2020, Section 7.1.3). When the learning rate decreases at an appropriate rate, and subject to relatively mild assumptions, stochastic gradient descent converges almost surely to local minimum (Bottou, 1998). However, there is theoretical evidence that in the deep learning context where the number of neural network parameters is very large, getting stuck in local minima is not a likely, and saddle points pose a greater challenge (see Dauphin et al., 2014). Adaptive gradient methods, such as the Adam optimizer we use in our work, address this challenge (Staib et al., 2019).

A.7.2 Passing redundant information to the equilibrium net. Heuristically, we found it helpful to pass redundant information to the neural network to stabilize the learning process. Since it increases the dimensionality of the input vector considerably, this is typically not done for methods like Smolyak sparse grids, or Gaussian processes. In our case, however, we found it helpful to increase the dimension of the input space by providing redundant information. Concretely, for the example studied in Section 5, we augmented the input state with the TFP shock $\eta(z)$, the depreciation $\delta(z)$, the aggregate capital K and labor L , the return on capital r and the wage for labor w , the aggregate production Y , the distribution of financial wealth across age groups $\mathbf{i}_{\text{fin}} = r \cdot \mathbf{k} + \mathbf{b} \cdot \min\{\kappa r, 1\}$, the distribution of labor income $\mathbf{i}_{\text{labor}} = w \cdot \mathbf{l}(z)$, the distribution of total wealth $\mathbf{i}_{\text{total}} = \mathbf{i}_{\text{labor}} + \mathbf{i}_{\text{fin}}$, and the transition probabilities of the exogenous shock $\Pi[z, :]$. To summarize, instead of passing the sufficient $2N + 1$ -dimensional vector $\mathbf{x} = [z, \mathbf{k}^T, \mathbf{b}^T]^T$ to the neural network, we pass a $4N + 12$ -dimensional vector $\mathbf{x}^* = [z, \eta(z), \delta(z), K, L, r, w, Y, \mathbf{k}^T, \mathbf{i}_{\text{fin}}^T, \mathbf{i}_{\text{labor}}^T, \mathbf{i}_{\text{total}}^T, \Pi[z, 1 : 4]]^T$ with redundant information to the neural network.⁴⁹

A.7.3 Dealing with infeasible predictions at the beginning of training. At the beginning of the training process, we initialize the parameters of the neural network with random values. Consequently, the approximated policy, which predicts random values, might predict savings that imply that some age groups consume negative amounts or that result in negative aggregate capital. Both of which cause the program to terminate with an error message. To deal with this issue, we replace infeasible values by feasible ones and add a punishment term to the loss function that trains the neural network not to predict policies that imply infeasible values. As learning proceeds and the approximate policies improve, these replacements are no longer necessary. Furthermore, it is important to verify that these replacements only occur at the beginning of the training process and never during simulation, after the training is finished. The following method is rather ad hoc, however, we found it to work sufficiently well:

- Set a punishment parameter: $\epsilon^{\text{punish}} = 10^{-5}$.
- If the predicted consumption is less than ϵ^{punish} , replace by $\epsilon^{\text{punish}} : \hat{c}_{\text{adjusted}}^i(\mathbf{x}) = \max(\hat{c}^i(\mathbf{x}), \epsilon^{\text{punish}})$.
- Add the punishment term $(\frac{1}{\epsilon^{\text{punish}}} \max(-\hat{c}^i(\mathbf{x}), 0))^2$ to the loss function.

That way, if the neural network predicts a policy implying negative consumption, the optimality conditions can still be evaluated. At the same time, the large punishment term will guide the parameter updates so that improved policies do not imply negative consumption.

A.8 An Example with an Analytical Solution. In this section, we evaluate the performance of our solution method in a setting in which an exact solution is known. The knowledge of an exact solution allows us to evaluate the precision of our approximate solution beyond the relative errors in the Euler equations, which we have used so far. Furthermore, we use this example to demonstrate that our method is applicable in settings in which approximate aggregation may not hold.

⁴⁹ Note that \mathbf{b} is implicitly passed to the neural network, because it is given by $\mathbf{b} = \frac{\mathbf{i}_{\text{fin}} - r \cdot \mathbf{k}}{\min\{\kappa r, 1\}}$.

We chose the same model as Krueger and Kubler (2004), which is an adapted version of the model in Huffman (1987). Although analytically solvable, this model is closely related to those we solve in this article.

A.8.1 The model. Next, for convenience and completeness, we outline this analytical test case. For readability, our notation here omits the dependence of variables on the node of the event tree, z^t , and we simply index them with time t , when there is no risk of confusion.

Households. Agents live for N periods and have log utility. Following Krueger and Kubler (2004), we assume that agents only work in the first period of their life: $l_t^s = 1$ for $s = 1$ and $l_t^s = 0$ otherwise. This implies that the aggregate labor supply is constant and equal to one: $L_t = 1$. Agents receive a competitive wage and can save in risky capital. Households cannot die with debt. Otherwise, there are no constraints or adjustment costs. The household's problem is given by

$$(A.44) \quad \max_{\{c_{t+i}^i, a_{t+i}^i\}_{i=0}^{N-1}} \mathbb{E}_t \left[\sum_{i=0}^{N-1} \log(c_{t+i}^i) \right]$$

subject to:

$$(A.45) \quad c_t^h + a_t^h = r_t k_t^h + l_t^h w_t,$$

$$(A.46) \quad k_{t+1}^{h+1} = a_t^h,$$

$$(A.47) \quad a_t^N \geq 0,$$

where c_t^h denotes the consumption of age group h at time t , a_t^h denotes the saving, k_t^h denotes the available capital in the beginning of the period, and r_t denotes the price of capital.

Firms. There is a single representative firm with Cobb–Douglas production function. The production function is given by

$$(A.48) \quad f(K_t, L_t, z_t) = \eta_t K_t^\alpha L_t^{1-\alpha} + K_t(1 - \delta_t),$$

where K_t denotes aggregate capital, L_t denotes the aggregate labor supply, α denotes the capital share in production, η_t denotes the stochastic TFP, and δ_t denotes the stochastic depreciation. The firm's optimization problem implies that the return on capital and the wage are given by $r_t = \alpha \eta_t K_t^{\alpha-1} L_t^{1-\alpha} + (1 - \delta_t)$ and $w_t = (1 - \alpha) \eta_t K_t^\alpha L_t^{-\alpha}$. Here, we study the special case where $L_t = l_t^1 = 1$.

A.8.2 Equilibrium. For completeness and convenience, we define the equilibrium we want to compute.

Competitive equilibrium.

DEFINITION A.3 (COMPETITIVE EQUILIBRIUM). A competitive equilibrium, given initial conditions $z_0, \{k_0^s\}_{s=1}^N$, is a collection of choices for households $\{(c_t^s, a_t^s)\}_{s=1}^N$ and for the representative firm $(K_t, L_t)_{t=0}^\infty$ as well as prices $(r_t, w_t)_{t=0}^\infty$, such that

1. given prices, households optimize,
2. given prices, the firm maximizes profits,
3. markets clear.

Functional rational expectations equilibrium. Here, a FREE is defined as follows:

DEFINITION A.4 (FUNCTIONAL RATIONAL EXPECTATIONS EQUILIBRIUM). A FREE consists of equilibrium functions $\theta_a : \mathcal{Z} \times \mathbb{R}^N \rightarrow \mathbb{R}^{N-1}$, where $\theta_a : \mathcal{Z} \times \mathbb{R}^N \rightarrow \mathbb{R}^{N-1}$ denotes the capital investment functions, such that for all states $\mathbf{x} := [z, \mathbf{k}^T]^T \in \mathcal{Z} \times \mathbb{R}^N$, where $z \in \mathcal{Z}$ denotes the exogenous shock and $\mathbf{k} = [k_1, \dots, k_N]^T$ denotes the endogenous state (i.e., the distribution of capital) with $k_1 = 0$, we have for all $i = 1, \dots, N - 1$:

$$(A.49) \quad u'(c^i(\mathbf{x})) = \beta E_z[r(\mathbf{x}_+)u'(\hat{c}^{i+1}(\hat{\mathbf{x}}_+))],$$

where $\mathbf{x}_+ = [z_+, 0, \theta_a(\mathbf{x})^T]^T$, where z_+ denotes the random exogenous shock in the next period, and where

$$(A.50) \quad r(\mathbf{x}) = f_K\left(\sum_{i=1}^N x_{1+i}, 1, x_1\right),$$

$$(A.51) \quad w(\mathbf{x}) = f_L\left(\sum_{i=1}^N x_{1+i}, 1, x_1\right),$$

$$(A.52) \quad c^i(\mathbf{x}) = \begin{cases} w(\mathbf{x}) - \theta_a(\mathbf{x})_i, & \text{for } i = 1, \\ r(\mathbf{x})x_{1+i} - \theta_a(\mathbf{x})_i, & \text{for } i = 2, \dots, N - 1, \\ r(\mathbf{x})x_{1+N} & \text{for } i = N. \end{cases}$$

A.8.3 Computing the equilibrium. We chose to present this model because it is closely related to the models we study in this article while an exact solution is available. In this section, we provide the exact solution and briefly reiterate how the equilibrium would be approximated using DEQNs.

Exact solution. Following Krueger and Kubler (2004), a FREE is given by

$$(A.53) \quad \theta_a(\mathbf{x}) = \beta \left[\frac{1 - \beta^{N-1}}{1 - \beta^N} w(\mathbf{x}), \frac{1 - \beta^{N-2}}{1 - \beta^{N-1}} r(\mathbf{x})k_2, \frac{1 - \beta^{N-3}}{1 - \beta^{N-2}} r(\mathbf{x})k_3, \dots, \frac{1 - \beta^1}{1 - \beta^2} r(\mathbf{x})k_{N-1} \right]^T.$$

This solution implies that aggregate capital evolves according to

$$(A.54) \quad K_t = \gamma_1 w(\mathbf{x}_{t-1}) + \gamma_2 w(\mathbf{x}_{t-2})r(\mathbf{x}_{t-1}) + \dots + \gamma_{N-1} w(\mathbf{x}_{t-(N-1)})r(\mathbf{x}_{t-1})r(\mathbf{x}_{t-2}) \dots r(\mathbf{x}_{t-(N-2)}),$$

where $\gamma_i := (\beta^i - \beta^N)/(1 - \beta^N)$.

Approximating the solution with DEQNs. We seek to find parameters ρ of a deep neural network \mathcal{N}_ρ , such that it approximates the true equilibrium policies:

$$(A.55) \quad \hat{\theta}_a(\mathbf{x}) := \mathcal{N}_\rho(\mathbf{x}) \approx \theta_a(\mathbf{x}).$$

Since, in general, the true policy $\theta_a(\mathbf{x})$ is unknown, we follow the algorithm described in this article and find parameters ρ by minimizing the errors in the equilibrium conditions with variants of mini-batch gradient descent. We can plug market clearing and the firms' optimization into the Euler equations so that the Euler equations characterize the recursive equilibrium. Given neural network parameters ρ , which imply approximate policy functions $\hat{\theta}_a(\mathbf{x})$, the errors in the equilibrium conditions at state \mathbf{x}_j are given by

$$(A.56) \quad e_{EE}^i(\mathbf{x}_j) := -u'(\hat{c}^i(\mathbf{x}_j)) + \beta E_z[r(\hat{\mathbf{x}}_+)u'(\hat{c}^{i+1}(\hat{\mathbf{x}}_+))], \text{ for } i = 1 \dots N - 1,$$

where

$$(A.57) \quad \hat{\mathbf{x}}_+ = \left[z_+, 0, \hat{\theta}_a(\mathbf{x})^T \right]^T,$$

and where z_+ denotes the random exogenous shock in the next period, and

$$(A.58) \quad r(\mathbf{x}) = f_K \left(\sum_{i=1}^N x_{1+i}, 1, x_1 \right),$$

$$(A.59) \quad w(\mathbf{x}) = f_L \left(\sum_{i=1}^N x_{1+i}, 1, x_1 \right),$$

$$(A.60) \quad \hat{c}^i(\mathbf{x}) = \begin{cases} w(\mathbf{x}) - \hat{\theta}_a(\mathbf{x})_i, & \text{for } i = 1, \\ r(\mathbf{x})x_{1+i} - \hat{\theta}_a(\mathbf{x})_i, & \text{for } i = 2, \dots, N-1, \\ r(\mathbf{x})x_{1+N} & \text{for } i = N. \end{cases}$$

The relative errors in the Euler equations are given by

$$(A.61) \quad e_{\text{REE}}^i(\mathbf{x}_j) := \frac{u'^{-1}(\beta E_{z_j}[r(\hat{\mathbf{x}}_{j,+})u'(\hat{c}^{i+1}(\hat{\mathbf{x}}_{j,+}))])}{\hat{c}^i(\mathbf{x}_j)} - 1$$

and the loss function for the DEQN is given by

$$(A.62) \quad \ell_{\mathcal{D}_{\text{train}}}(\rho) := \frac{1}{|\mathcal{D}_{\text{train}}|} \frac{1}{N-1} \sum_{\mathbf{x}_j \in \mathcal{D}_{\text{train}}} \sum_{i=1}^{N-1} e_{\text{REE}}^i(\mathbf{x}_j)^2,$$

where $\mathcal{D}_{\text{train}}$ denotes a set of states collected by simulating the economy using the current approximation of the policy functions.

A.8.4 Parameterization and hyperparameters. We solve the model for $N = 6$ agents and base our parameterization on Krueger and Kubler (2004). The chosen economic parameters as well as the chosen hyperparameters are given in Table A.8.

A.8.5 Assessing the quality of the solution. We train the neural network for a total of 10,000 episodes. We first evaluate the quality of the obtained policy functions by looking at the relative errors in the Euler equations, since this is the measure we focus on in the settings we are interested in and where no exact solution is available. Table A.9 shows the relative errors in the Euler equations on 15,000 randomly simulated states.⁵⁰

The mean errors are in the order of $\sim 10^{-4}$, whereas the 99.9th percentile is below 1%. Since this is a much simpler model than our benchmark model, the errors are slightly lower than the errors we obtained for the models discussed in the previous sections.

Since the exact solution is available in this model, we now compare the approximate policy learned by our method to the precise solution. Figure A.8 shows the savings decisions learned by the DEQN (shown as stars) as well as the precise solution (shown as circles) as a function of the agents' capital at the beginning of each period in a scatter plot. The figure shows only 200 simulated states for better readability. The colors indicate the four exogenous shocks. Visually, the policies are indistinguishable. Table A.10 shows statistics of the errors in the learned savings decisions.

⁵⁰ We simulate 16,000 periods and throw away the first 1,000.

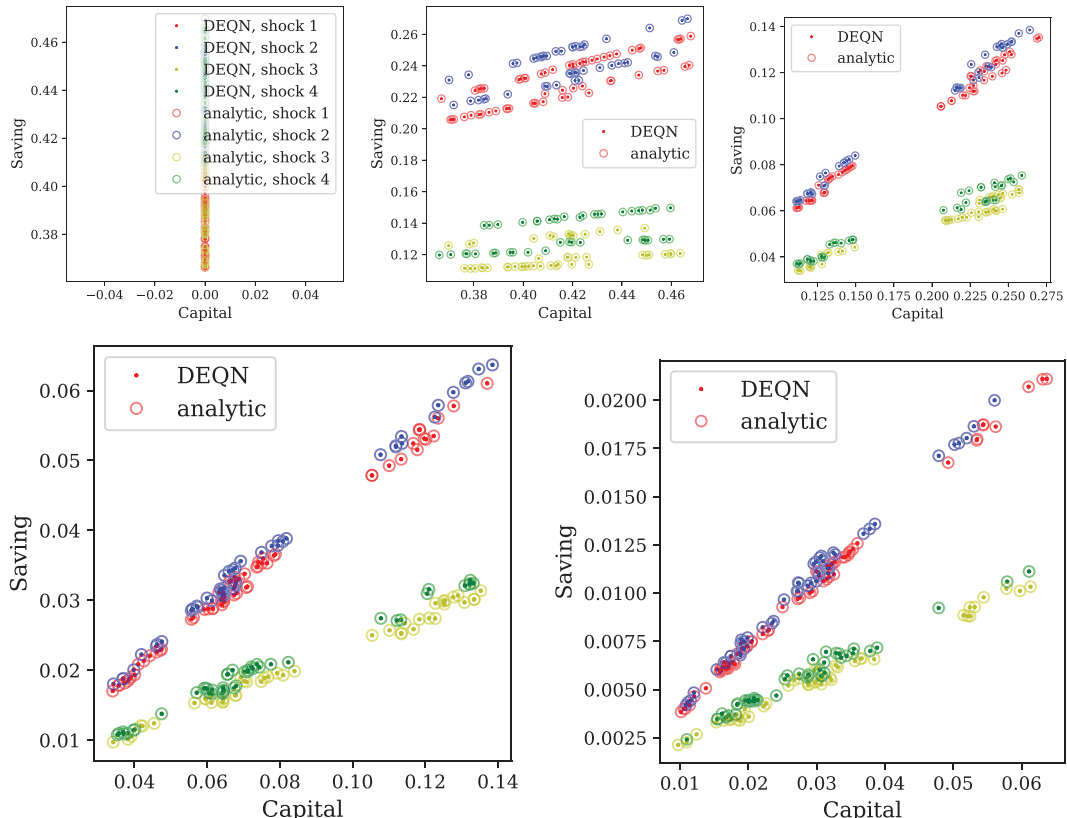
TABLE A.8
PARAMETERIZATION OF THE ECONOMIC PARAMETERS (TOP) AND THE HYPERPARAMETERS OF THE NEURAL NETWORK (BOTTOM)

	Discount Factor β	Relative Risk Aversion γ	Capital Share α	TFP η	Depreciation δ	Persistence TFP $P(\eta_{t+1} = 1.05 \eta_t = 1.05)$	Persistence Depreciation $P(\delta_{t+1} = 0.5 \delta_t = 0.5) P(\delta_{t+1} = 0.5 \delta_t = 0.9)$
Number Groups N	6	0.7	1	0.3	{0.95, 1.05}	{0.5, 0.9}	{0.5, 0.9}
Learning Rate α_{learn}	0.00001	Periods per Episode T	Epochs per Episode N^{epochs}	Mini-Batch Size m	Nodes Hidden Layers	Activations Hidden Layers	Activations Hidden Layers
		12,800	20	640	100 50	relu relu	relu relu

TABLE A.9
RELATIVE EULER EQUATION ERRORS ON 15,000 SIMULATED PERIODS

	Mean	Max	0.1	10	50	90	99.9
Rel Ee capital [log10]	-3.4	-2.4	-6.4	-4.4	-3.6	-3.0	-2.5

NOTE: The columns show the mean and the max errors as well as the 0.1st, 10th, 50th, 90th, and 99.9th percentile.



NOTES: Exact and approximate saving decisions plotted against capital holding at the beginning of the period for 200 simulated states for each of the 6 age groups. The circles show the exact solution, and the stars show the solutions learned by the deep equilibrium net after training for 10,000 episodes with parameters shown in Table A.8. The first row shows the policies for age groups 1, 2, and 3, the second row shows the policies for age groups 4 and 5. The policy of the sixth age group is trivial (consume everything and save nothing). Shock 1 is denoted in red, shock 2 in blue, shock 3 in yellow, and shock 4 in green. The TFP and depreciation (η, δ) of shock 1, shock 2, shock 3, and shock 4 are $(0.95, 0.5)$, $(1.05, 0.5)$, $(0.95, 0.9)$, and $(1.05, 0.9)$, respectively.

FIGURE A.8

EXACT AND APPROXIMATE SAVING DECISIONS

The mean errors in the policy functions are of order $\sim 10^{-4}$ and the maximum errors stay below 0.2%. In the next section, we study the extent to which an accumulation of errors could influence aggregates in this model.

A.8.6 Approximate aggregation. This section serves two purposes. First, we want a measure of the accumulation of errors with time as we simulate the economy forward. Second, we want to see to what extent approximate aggregation, as in the seminal work of Krusell and Smith (1998), holds in this model.

Following Krueger and Kubler (2004), we only use the log of aggregate capital to forecast next-period's capital contingent on the exogenous shock when assessing the approximate

TABLE A.10
STATISTICS OF THE RELATIVE ERRORS IN THE LEARNED POLICY FUNCTIONS ON 15,000 SIMULATED PERIODS

	Mean	Max	0.1	50	99.9
Relative policy errors age group 1 [%]	0.03	0.14	0.00	0.03	0.13
Relative policy errors age group 2 [%]	0.02	0.09	0.00	0.01	0.08
Relative policy errors age group 3 [%]	0.02	0.10	0.00	0.02	0.09
Relative policy errors age group 4 [%]	0.01	0.05	0.00	0.01	0.14
Relative policy errors age group 5 [%]	0.01	0.06	0.00	0.01	0.04
Relative policy errors age group 1 [log10]	-3.47	-2.85	-6.08	-3.54	-2.88
Relative policy errors age group 2 [log10]	-3.82	-3.06	-6.72	-3.91	-3.10
Relative policy errors age group 3 [log10]	-3.69	-2.99	-6.40	-3.75	-3.04
Relative policy errors age group 4 [log10]	-4.09	-3.29	-7.11	-4.26	-3.37
Relative policy errors age group 5 [log10]	-3.92	-3.33	-6.70	-4.00	-3.35

NOTE: The columns show the mean and the max errors as well as the 0.1st, 50th, and 99.9th percentile.

TABLE A.11
STATISTICS OF THE RELATIVE ERRORS IN THE SEQUENCE OF AGGREGATE CAPITAL COMPARED TO THE EXACT SOLUTION WHEN SIMULATING 15,000 TIME PERIODS

	Mean	Max
DEQN [%]	0.019	0.13
Log-linear forecast [%]	0.24	1.3
DEQN [log10]	-3.72	-2.88
Log-linear forecast [log10]	-2.62	-1.87

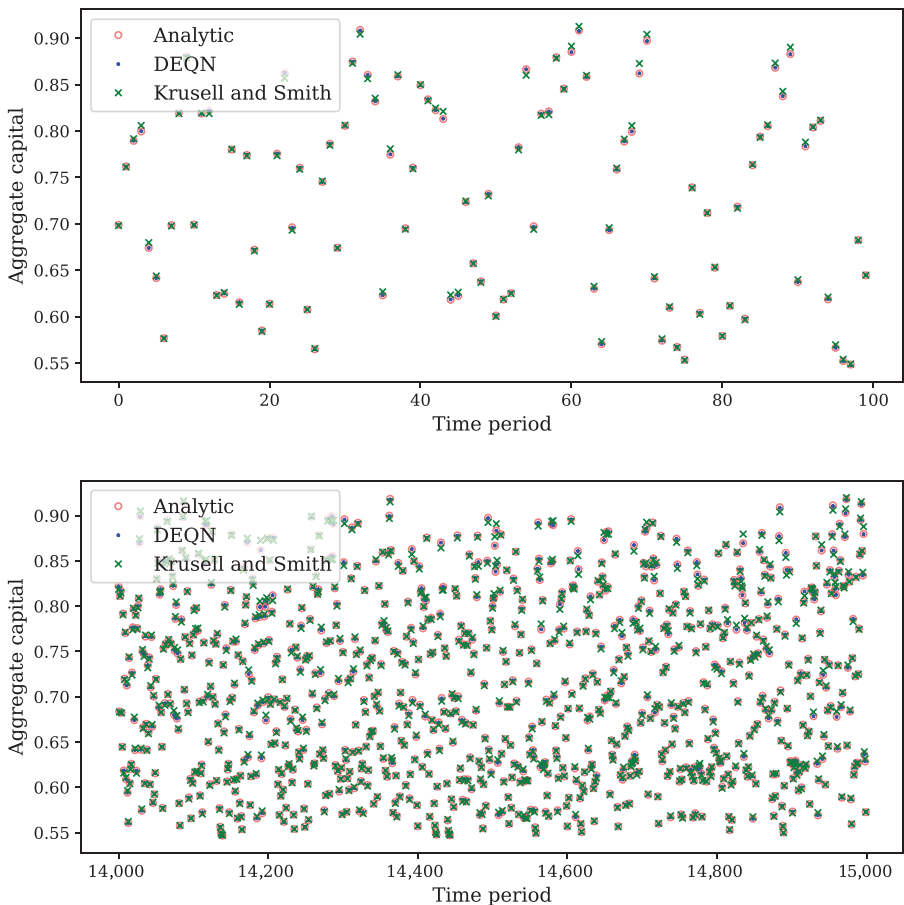
aggregation in this model.⁵¹ More precisely, the functional form of the forecasting rule is given by

$$(A.63) \quad \log(K_{t+1}) = \phi_{0,z} + \phi_{1,z} \log(K_t),$$

where we estimate the parameters $\phi_{0,z}$ and $\phi_{1,z}$ with an ordinary least squares regression for each of the four shocks. The R^2 values of the corresponding regressions are between 0.9978 and 0.9979. The corresponding mean forecasting errors are between 0.16% and 0.24%. The corresponding maximum forecasting errors are between 0.61% and 0.98%.

Following Den Haan (2010) and Winberry (2018), we also analyze the extent to which approximate aggregation holds by simulating the linear forecast of aggregate capital forward without updating the value of aggregate capital. We do the same using the solution learned by the DEQN and compare the sequences of aggregate capital on 15,000 simulated states. Table A.11 reports the mean and maximum of the relative error of the sequence of aggregate capital compared to the exact solution. The mean relative errors in the sequence of aggregate capital are more than one order of magnitude lower using the policies learned by the DEQN than when using a log-linear approximation of the law of motion. To illustrate this point, Figure A.9 shows the simulated values of aggregate capital for the exact solution (red circles), using the policies learned by the DEQNs (blue dots), and when using a log-linear forecast (green crosses). For readability, we show the first 100 and the last 1,000 periods. The figure shows that the evolution of aggregate capital obtained from the policies learned by the DEQN is visually indistinguishable from the exact solution, whereas the evolution using a log-linear forecast shows visible differences. This further illustrates the point that our method applies to settings in which approximate aggregation does not obtain. For both approximate solutions, we find no evidence that the quality of the solution deteriorates when simulating many periods.

⁵¹ The regressions use 15,000 simulated states in total, approximately 3,750 states for each shock.



NOTES: The red circles show the exact evolution, the blue stars show the evolution implied by the policies learned by the deep equilibrium net, and the green crosses show the evolution when using a log-linear forecast. The upper figure shows the first 100 and the lower figure the last 1,000 periods of 15,000 simulated periods.

FIGURE A.9

EXACT AND APPROXIMATED EVOLUTION OF AGGREGATE CAPITAL

As noted in Krueger and Kubler (2004), in this simple model, errors in forecasting the aggregate capital stock do not translate to welfare losses for the agents. This is because, in the simple model presented here, the optimal decision does not depend on interest rates.

A.9 Parallelization Scheme. To speed up the solution process of the computational method introduced in this article by orders of magnitude, we present a generic and highly-scalable parallelization scheme that allows us to make efficient use of state-of-the-art high-performance computing (HPC) facilities, whose performance nowadays can reach up to hundreds of Petaflop/s (see, e.g., <https://www.top500.org>). The use of modern HPC hardware may enable us to solve hard problems with many state variables in a relatively short time and to tackle problems that have, thus far, been out of reach.

Our algorithm can be described as iterating on two phases: the simulation phase, where new training data are generated using the previous iteration's neural network, and the learning phase, where the neural network is trained on the new data. We make use of contemporary frameworks to parallelize both phases. More precisely, our method is parallelized by combining TensorFlow (Abadi et al., 2015) with Horovod (Sergeev and Del Balso, 2018).

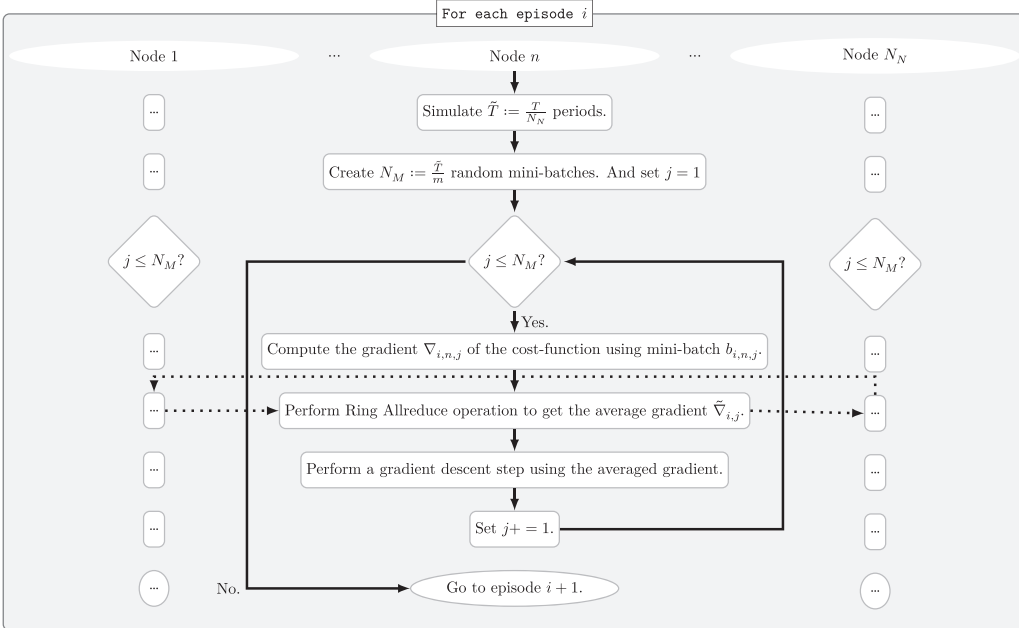


FIGURE A.10

SCHEMATIC ILLUSTRATION OF THE PARALLELIZATION SCHEME

Horovod is an open-source distributed training framework for TensorFlow (and other deep learning codes) that was developed to make distributed deep learning fast and easy to use. The core idea behind distributed deep learning is the following: first, training data are distributed across the compute nodes in a cluster. On each node, the gradient of the cost function with respect to the parameters of the neural network on its given batch of data is determined. Finally, all nodes are synchronized such that the gradient descent step can be taken using the average gradient across the nodes. Horovod implements the synchronization and calculation of the average gradient using a Ring Allreduce operation.^{52,53}

In addition, our simulation phase also lends itself nicely to parallelism: each node simulates the part of the training data, which it will then use to compute the gradient during the learning phase. This also limits the communication between the nodes. The only communication required among the nodes is the communication of the gradients for the Ring Allreduce operation. In Section A.9.1, we provide additional details on the parallelization and in Section A.9.2, we demonstrate that this parallelization scheme can accelerate the computations by orders of magnitude.

A.9.1 Hybrid parallelization scheme for heterogeneous HPC systems. In the following, we detail the hybrid parallelization scheme, which is schematically displayed in Figure A.10. Let N_N denote the number of nodes available in the computing system and $n \in \{1, \dots, N_N\}$ denote a specific node. Each node has a copy of the same neural network with identical weights, such that $\rho_i = \rho_j, \forall i, j \in \{1, \dots, N_N\}$. Then, each node has a different seed for random number generators to ensure that each node generates different data during the simulation phase. The following summarizes the parallelization scheme for a given episode i :

⁵² The Horovod source code was based on the Baidu TensorFlow-allreduce repository written by Andrew Gibiansky and Joel Hestness (cf, <https://github.com/baidu-research/tensorflow-allreduce>).

⁵³ Ring Allreduce is an efficient version of the Allreduce operator commonly used in HPC. Ring Allreduce mitigates a bottleneck in the amount of data sent between nodes. Allreduce is tied to another operator, in our case, the mean. At the end of Ring Allreduce, each node has a copy of the Allreduced data.

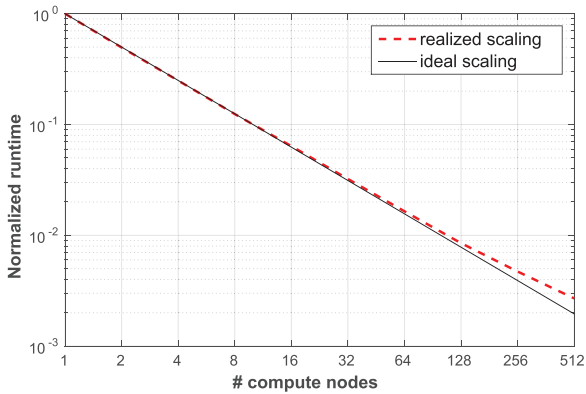


FIGURE A.11

STRONG SCALING

Simulation phase

On each node n :

- Draw a random sequence of exogenous shocks and simulate $\tilde{T} := T/N_N$ periods using the policy encoded by the neural network \mathcal{N}_{ρ_n} .
- Create $N_m := \tilde{T}/m$ random mini-batches. We denote the mini-batches as $b_{i,n,j}$ and the set of mini-batches by $\mathcal{B}_{i,n} = \cup_{j=1}^{N_m} b_{i,n,j}$.

Learning phase

On each node n :

For each mini-batch $b_{i,n,j} \in \mathcal{B}_{i,n}$ (i.e., for $j = 1, \dots, N_m$):

- Compute the gradient of the cost function with respect to the parameters of the neural network ρ_n , denoted as $\nabla_{i,n,j}$.
- Perform a Ring Allreduce operation to obtain the average gradient across nodes,

$$\tilde{\nabla}_{i,j} = \frac{1}{N_N} \sum_{k=1}^{N_N} \nabla_{i,k,j}.$$

- Update the parameters of the neural network doing a gradient descent step as implied by the average gradient ($\tilde{\nabla}_{i,j}$). New weights: $\rho_n^{\text{new}} = \rho_n^{\text{old}} - \alpha^{\text{learn}} \tilde{\nabla}_{i,j}$.

A.9.2 Strong scaling. In this section, we report the strong scaling and the efficiency of our code on a simple benchmark problem, similar to the model studied in Section 5. This scaling test was carried out on the Cray XC50 “Piz Daint” system that is installed at the Swiss National Supercomputer Centre (CSCS). Cray XC50 compute nodes combine Intel Xeon E5-2690 v3 CPUs (12 cores, 64 GB RAM) with one NVIDIA P100 GPU that is equipped with 16 GB of memory.

In our strong scaling example, we chose the network architecture to be 1,500 nodes for the two hidden layers, a batch size of 150,000, and an episode length of 76,800,000 periods. These numbers are larger than those used above, reflecting the fact that we aim at using larger networks and batch sizes to study more complicated models in the future. Figure A.11 shows the resulting strong scaling and efficiency. The runtime is normalized by the time it takes a single node to simulate and train on one episode. As displayed, we find excellent strong scaling properties, with a parallel efficiency above 75%, even at 512 hybrid compute nodes.

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