END-TO-END LARGE PORTFOLIO OPTIMIZATION FOR VARIANCE MINIMIZATION WITH NEURAL NETWORKS THROUGH COVARIANCE CLEANING

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ABSTRACT

We develop a rotation-invariant neural network that provides the global minimum-variance portfolio by jointly learning how to lag-transform historical returns and how to regularise both the eigenvalues and the marginal volatilities of large equity covariance matrices. This explicit mathematical mapping offers clear interpretability of each module's role, so the model cannot be regarded as a pure blackbox. The architecture mirrors the analytical form of the global minimum-variance solution yet remains agnostic to dimension, so a single model can be calibrated on panels of a few hundred stocks and applied, without retraining, to one thousand US equities—a cross-sectional jump that demonstrates robust out-of-sample generalisation. The loss function is the future realized minimum portfolio variance and is optimized end-to-end on real daily returns. In out-of-sample tests from January 2000 to December 2024 the estimator delivers systematically lower realised volatility, smaller maximum drawdowns, and higher Sharpe ratios than the best analytical competitors, including stateof-the-art non-linear shrinkage. Furthermore, although the model is trained end-to-end to produce an unconstrained (long-short) minimum-variance portfolio, we show that its learned covariance representation can be used in general optimizers under long-only constraints with virtually no loss in its performance advantage over competing estimators. These gains persist when the strategy is executed under a highly realistic implementation framework that models market orders at the auctions, empirical slippage, exchange fees, and financing charges for leverage, and they remain stable during episodes of acute market stress.

Keywords Covariance cleaning · Portfolio optimization · Neural Network · Global Minimum Variance · End-to-End

1 Introduction

Portfolio management has been fundamental to financial practice since the earliest securities markets emerged. Following Markowitz's introduction of Modern Portfolio Theory (MPT) [1], academic and industry efforts have prioritized the construction of portfolios that optimize risk-adjusted returns. This framework employs historical return data and estimates of pairwise asset dependencies, typically captured by covariance matrices, to determine asset weightings.

Classic portfolio optimization proceeds in two logical steps: parameter estimation followed by the optimization itself. The optimization phase depends on forecasts of asset return vectors and their covariance matrix. Predicting these quantities is notoriously difficult, and a vast literature seeks either to improve forecast accuracy or to develop stochastic frameworks that explicitly accommodate their uncertainty. Forecasting returns is especially challenging because it reflects the degree of market efficiency observed in financial markets [2]. Estimating covariances likewise involves fundamental statistical estimation issues [3] and must account for their time-varying nature. Nonetheless, covariance forecasting is relatively more tractable, since improvements in the covariance estimate cannot be directly translated into profit in simple trading strategies without resort to complex financial instruments.

The recent development of powerful machine learning approaches suggests that their use in portfolio optimization can provide results compatible with or better than state-of-the-art academic and professional solutions formalized in terms of mathematical finance models. Machine learning approaches to portfolio optimization can present different schemes. The most widespread approaches are: a) predict-then-optimize, b) decision-focused learning, and c) end-to-end approach [4]. Recent studies illustrate the extent of Neural Network (NN) applications to portfolio optimisation, yet they also expose recurring limitations that motivate the present work. Mesquita et al. [5] couple a Heterogeneous AutoRegressive filter with a multilayer perceptron to obtain positive-definite covariance forecasts, but the experiment involves only five equities and omits a Global Minimum Variance (GMV) benchmark, so scalability and incremental value remain unclear. Burda and Schroeder [6] embed a GO-GARCH model within a Long-Short Term Memory (LSTM) network to generate weights for the 100 largest U.S. stocks; the design yields modest gains over equal-weight and DCC strategies and is susceptible to survivor bias because delisted firms are excluded ex post. Huang et al. [7] adopt a two-stage network that first screens and then weights equities, yet the screening step discards securities that delist during the backtest, thus leaking future information. Reis et al. [8] combine a CNN and an LSTM to forecast covariances for a 14-asset mix of equities and fixed-income ETFs; fixed-income dominance in the GMV solution limits the interpretability of the results, and further evaluation on larger universes or on purely equity portfolios would be necessary to assess the model's broader applicability. Collectively, these contributions demonstrate the promise of machine learning but highlight four issues: small or static universes, sub-optimal or heterogeneous benchmarks, potential data leakage, and uncertain behaviour when the cross-sectional dimension approaches the sample length. Addressing these gaps is the central objective of the architecture introduced below.

The present study introduces an NN architecture that implements an end-to-end GMV portfolio optimizer by mapping each analytical step of the classical solution into a modular network. Historical returns for n assets over Δt time periods are first processed through a learnable lag-transformation module that applies distinct weights and soft-clipping parameters across look-back lags. The transformed returns then feed into a bidirectional LSTM-based correlation denoising module that adapts to the aspect ratio $q=n/\Delta t$, and into a parallel feed-forward network that estimates inverse marginal volatilities. All modules are calibrated jointly to minimize the Out-Of-Sample (OOS) portfolio variance in an end-to-end fashion. The correlation denoising module represents the greatest learning challenge, since its multivariate nature and the inherent symmetry of correlation matrices impose complex constraints on the network. If the architecture is correctly designed, calibration on datasets with n in a narrow range should yield a parameter set that generalizes to different values of n. This expectation is borne out in our experiments. OOS tests on a universe of one thousand U.S. equities covering January 2000 through December 2024, using calibration on just a few hundred stocks, confirm that our architecture delivers lower realised variance and higher Sharpe ratios than state-of-the-art methods. When evaluated under realistic trading procedures and cost assumptions, it also produces consistently smaller drawdowns.

A key aspect of our model is associated with the determination of the covariance matrix. When estimated from finite samples, empirical covariance matrices tend to be ill-conditioned and sample covariance presents estimation errors [3, 9, 10], often resulting in suboptimal portfolio allocations. A widely adopted remedy is to apply Rotation Invariant Estimators (RIEs), which adjust the sample eigenvalues while retaining the sample eigenvectors [11]. Within this framework, optimality is measured by the Mean Squared Error (MSE), equivalent to the Frobenius norm distance between the estimator and the true covariance matrix. The theoretical optimum is the so-called Oracle Estimator (OE), defined as the covariance matrix that minimizes this Frobenius distance. In practice, however, the OE cannot be realized, since it requires prior knowledge of the unknown true covariance.

One of the earliest regularization techniques is eigenvalue clipping. Laloux et al. [9] showed that the bulk of the eigenvalue spectrum of empirical financial covariance matrices closely follows the Marchenko–Pastur distribution for random noise. Eigenvalue clipping therefore replaces all sample eigenvalues that lie within this noise bulk by their average value. Later, Ledoit and Wolf [10] introduced the Linear Shrinkage (LS) estimator, which constructs a convex combination of the sample covariance matrix and a diagonal target matrix. This estimator remains well–defined when the number of assets n exceeds the number of observations Δt . Its optimality in minimizing MSE holds strictly under the assumption that the true covariance matrix is drawn from an inverse–Wishart distribution [12]. In the last decade, Ledoit and collaborators introduced the NonLinear Shrinkage (NLS) estimator as an optimal method for covariance

regularization [13]. Deriving the analytical form of the Ledoit–Péché (LP) estimator is challenging because it requires detailed knowledge of the sample eigenvector distribution [14]. The Quantized Eigenvalues Sampling Transform (QuEST) delivers the most accurate numerical implementation of the LP estimator by discretizing the limiting spectral distribution [15], but it incurs substantial computational cost. To alleviate this burden, the direct kernel estimator [16] and the inverse–Wishart regularization approach [11] offer closed-form approximations that closely mimic the LP solution. More recently, Ledoit and Wolf proposed the Quadratic-Inverse Shrinkage (QIS) estimator [17], which is considered the state-of-the-art technique in financial covariance filtering by combining computational efficiency with high estimation accuracy.

Although the theoretical framework for covariance filtering is well established, its practical implementation remains challenging due to the pronounced non-stationarity of financial covariances. Market regimes evolve continuously under shifting economic conditions and investor behavior, so a covariance matrix estimated from past returns may poorly reflect current asset dependencies. Moreover, NLS theory assumes that sampling noise is the sole source of discrepancy between the sample and true covariance matrices and relies on the thermodynamic limit in which the number of assets n and observations Δt both tend to infinity with fixed aspect ratio $q = n/\Delta t$. In practice, extending Δt to satisfy this limit only exacerbates non-stationarity, as longer historical windows encompass multiple structural shifts. Finally, the optimality of the LP estimator depends on the existence of finite twelfth-order moments of asset returns [14], yet empirical evidence indicates that equity returns exhibit heavy tails with tail exponents near four, rendering higher-order moments effectively unbounded. Methods based on k-fold Cross-Validation (CV) have been proposed to relax the stringent assumptions of nonlinear shrinkage theory [18, 12, 19]. These CV estimators achieve accuracy comparable to more complex schemes while permitting straightforward adaptations to specific data features. For example, Tan et al. [20] introduced an exponential decay factor into the return series within the CV framework. By applying exponential weights, one can maintain a long calibration window—approaching the thermodynamic limit—while emphasizing the most recent observations. This modification partially resolves the variance-bias trade-off caused by non-stationarity and structural shifts in financial covariances.

An alternative to RIEs employs hierarchical clustering to filter the empirical correlation matrix [21, 22, 23]. Unlike RIEs, clustering-based filters modify both eigenvalues and eigenvectors by aggregating assets into empirically determined groups. This approach exploits the fact that financial assets are often organized into a clear hierarchy of sectors and sub-sectors [24]. Although it lacks full theoretical support, clustering filters have, in some instances, yielded superior out-of-sample portfolio performance compared with RIEs [25, 26, 22, 23, 27]. By enforcing the observed hierarchical relationships, these methods produce the most parsimonious covariance approximation that exactly reproduces the data's cluster structure.

The Engle Dynamical Conditional Covariance (DCC) model [28] represents a principled attempt to accommodate non-stationarity by modeling time-varying correlations through GARCH-type dynamics. It specifies separate equations for the conditional variances and then characterizes the evolving correlations of the standardized returns via a time-varying correlation matrix. In principle, this approach should adapt to shifting market regimes and capture persistence in asset comovements. In practice, however, DCC is computationally intensive, often suffers from convergence issues, and yields OOS portfolios whose risk—return profiles rarely surpass those obtained with simpler static filters [29]. In contrast, the Average Oracle (AO) method [30, 29] abandons time-varying adjustments altogether. It fixes a set of eigenvalues calibrated over an extended historical window and applies them uniformly to all subsequent covariance estimates. This static correction ignores short-term noise but captures the enduring structure of the market. Empirical tests show that AO consistently outperforms both DCC-based and other NLS techniques, suggesting that, under strong non-stationarity, long-term averages provide more reliable covariance estimates than complex dynamic models [29, 31].

The AO method effectively serves as a zero-order baseline, applying fixed eigenvalue corrections without any attempt to anticipate future shifts. An ideal covariance estimator, by contrast, would move beyond denoising past returns and actively predict forthcoming covariance dynamics. NNs naturally suggest themselves for this role, since they can model highly nonlinear relationships between input data and desired outputs. Two main strategies can be used when training an NN for GMV optimization. The first is based on learning a covariance matrix estimator, which is then used to compute the portfolio weights with a Quadratic Programming (QP) minimization. This approach has the advantage of being versatile and adaptable to different minimization problems, especially if they are convex and optimally solvable. However, it has the drawback of requiring the estimation of a covariance matrix, which needs to be positive definite. In addition, the loss function should be taken as some type of distance between the predicted covariance matrix and the population covariance matrix, usually the Frobenius distance. This presents a challenge, as the Frobenius distance is not a good metric for covariance matrices in practical portfolio optimization problems [32, 33]. The second approach is to train an NN to predict portfolio weights directly. This method might allow for a straightforward optimization of the desired loss function if the NN architecture is designed in a tailored way. Depending on the architecture, the NN can internally store a representation of the covariance matrix or some simplified version. In this paper, we adopt the latter approach.

At the heart of our architecture lies the correlation-denoising module (Sec. 3.2), which implements a RIE of the correlation matrix. By applying a RIE, one forces all permutational symmetries already encoded in its mathematical formulation. To perform this eigenvalue filtering efficiently, the symmetry of the eigenvalue transformation itself must be respected. We therefore take inspiration from the well-known mapping of sample eigenvalues to a one-dimensional Coulomb gas [34], in which each eigenvalue behaves as repelling identical charges confined by the population spectrum. In our implementation, the ordered sequence of sample eigenvalues is processed through a bidirectional LSTM that enforces permutation equivariance and captures the local interactions among neighboring "charges". The network outputs a q-dependent NLS function that adjusts each sample eigenvalue while leaving the empirical eigenvectors intact. Two auxiliary modules prepare the raw return data for this denoising step. A learnable lag-transformation network weights look-back lags and applies soft clipping to control temporal influence and outliers (Sec. 3.1). A marginal-volatility network converts each asset's sample standard deviation into an inverse volatility scale (Sec. 3.3). These modules work in concert to present the LSTM with input representations whose temporal dependencies and scales are optimally calibrated. Finally, the cleaned eigenvalues recombine with the inverse marginal volatilities to form an estimated inverse covariance matrix, from which GMV portfolio weights follow in closed form. Calibrating all modules jointly to minimize realized OOS variance enables the model to predict future covariance dynamics under non-stationary market regimes. This end-to-end design thus overcomes the limits of Frobenius-norm-based shrinkage and static filters and delivers a dynamically adaptive estimator of portfolio risk.

2 Mathematical Background

2.1 GMV Portfolios under Finite-Sample Risk-Mode Uncertainty

Consider an idealised setting in which an investor observes $\Delta t_{\rm in} > n$ independent realisations of the n-dimensional return vector $\boldsymbol{r}_t = \{r_{t1}, r_{t2}, \dots, r_{tn}\}^{\top}$. We refer to these observations as the In-Sample (IS) data and assume each \boldsymbol{r}_t is drawn from a multivariate normal distribution with zero mean and covariance matrix $\boldsymbol{\Sigma}$. The task of the GMV portfolio is to find weights $\boldsymbol{w} = \{w_1, w_2, \dots, w_n\}^{\top}$ that minimise portfolio variance subject to the full-investment constraint $\mathbf{1}^{\top} \boldsymbol{w} = 1$. Replacing the unknown covariance matrix with its maximum-likelihood estimator $\hat{\boldsymbol{\Sigma}} = \boldsymbol{\Sigma}_{\rm MLE}$ yields the typical application of the GMV portfolio

$$w = \frac{\widehat{\Sigma}^{-1} \mathbf{1}}{\mathbf{1}^{\top} \widehat{\Sigma}^{-1} \mathbf{1}}.$$
 (1)

Performance is later assessed on an independent OOS period of length $\Delta t_{\rm out}$, sampled from the same distribution $\mathcal{N}(\mathbf{0}, \mathbf{\Sigma})$.

Sampling error inflates the realised risk of the GMV portfolio $\sigma_{\text{PTF}}^2 = \boldsymbol{w}^{\top} \boldsymbol{\Sigma}_{\text{out}} \boldsymbol{w}$. Under multivariate normality, the expected OOS variance is

$$\mathbb{E}\left[\sigma_{\text{PTF}}^{2}\right] \approx \frac{1}{\mathbf{1}^{\top} \mathbf{\Sigma}^{-1} \mathbf{1}} \left(1 + \frac{n}{\Delta t_{\text{in}} - n}\right) = \sigma_{\star}^{2} \left(1 + \frac{q_{\text{in}}}{1 - q_{\text{in}}}\right),\tag{2}$$

where $q_{\rm in}=n/\Delta t_{\rm in}$ and $\sigma_{\star}^2=(\mathbf{1}^{\top}\mathbf{\Sigma}^{-1}\mathbf{1})^{-1}$ denotes the population GMV portfolio variance, i.e., an idealized portfolio that is built and tested on the population covariance $\mathbf{\Sigma}$ [35]. Note that the inflation factor depends only on the estimation window, and it diverges when $\Delta t_{\rm in} \to n$. Interestingly, $\Delta t_{\rm out}$ affects only the dispersion of the variance estimator.

Since most covariance-cleaning procedures act through spectral modifications of the MLE correlation or covariance matrices, it is instructive to express the GMV portfolio in the eigenbasis and examine the contribution of each risk mode.

Let $\Sigma = UZU^{\top}$ with eigenvalues ζ_k and eigenvectors u_k . The GMV portfolio can be decomposed as

$$\boldsymbol{w} \propto \sum_{k=1}^{n} \frac{c_k}{\zeta_k} \boldsymbol{u}_k, \qquad c_k := \boldsymbol{u}_k^{\top} \mathbf{1} = \sum_{i=1}^{n} u_{ik}.$$
 (3)

We orient each eigenvector so that $c_k \geq 0$; changing the sign of an eigenvector leaves Σ unchanged, so this choice is harmless. The coefficient c_k gauges how closely the eigen-portfolio u_k points toward the budget line 1. If $c_k = 0$ the mode is orthogonal to 1 and has equal long and short exposure, hence no net investment. Allocating wealth to such a direction therefore increases variance without helping to deploy funds, so the GMV portfolio gives preference to modes with $c_k > 0$. Eq. (3) makes this explicit: eigen-portfolio u_k enters with weight c_k/ζ_k , the product of its alignment with the budget and the inverse of its own variance $\zeta_k = u_k^T \Sigma u_k$. Combining modes in this way spreads risk across orthogonal directions and achieves a total variance lower than that of any single eigen-portfolio.

Similarly, the eigendecomposition $C = V \Lambda V^{\top}$ of the correlation matrix leads to

$$w_i \propto \sum_{k=1}^n \frac{c_k}{\lambda_k} \frac{v_{ik}}{\sigma_i}, \qquad c_k := \sum_{i=1}^n \frac{v_{ik}}{\sigma_i}, \qquad i = 1, \dots, n;$$
 (4)

where σ_i is the marginal volatility of the asset *i*. The scaling by σ_i magnifies the influence of low-volatility assets and attenuates that of volatile ones. Although the resulting eigen-portfolios are not more orthonormal, they are still orthogonal; therefore, the interpretation of c_k and λ_k^{-1} remains unchanged.

In practice, one does not observe the true eigenbasis but only its sample estimate from finite data. RMT results indicate that when a population eigenvalue λ_k is well separated from the rest of the spectrum, the corresponding sample eigenvector \hat{v}_k tends to have a high overlap with the true v_k [36]. Unfortunately, such spectral isolation occurs primarily for the largest eigenvalues, which enter the global minimum-variance portfolio with weights proportional to $1/\lambda_k$ and thus contribute little to risk reduction. The smaller eigenvalues are tightly clustered in the bulk of the spectrum, causing their sample eigenvectors to mix and lose any precise orientation. As a result, estimated eigen-portfolios associated with these directions are dominated by sampling noise rather than true risk modes.

Furthermore, empirically realised OOS portfolio variances often exceed the theoretical benchmark by an order of magnitude [37, 26]. Two features of equity data might explain most of this gap. First, returns are heteroskedastic and exhibit time-varying correlations, so that $\Sigma_{\rm in}$ may differ substantially from $\Sigma_{\rm out}$ [38, 39, 40]. Second, return distributions are heavy-tailed, well-approximated by a Student-t law with four to six degrees of freedom [41, 42]. Because finite fourth moments are absent or unstable, the classical Wishart assumptions underlying RMT break down, amplifying estimation error [43].

If short selling is forbidden, one adds the element-wise constraint $w_i \ge 0$ to the GMV optimization. The problem remains convex but no longer has a closed-form solution; however, it can still be optimally solved by numerical quadratic programming. Empirically, it was observed in Ref. [26, 44] that long-only portfolios on the ML covariance estimator tend to perform similarly to cleaned covariance estimators.

2.2 Eigenvalue Cleaning for MSE Minimization

Having established the problem and its connection to the portfolio's eigen-risk modes, we now outline how conventional covariance-cleaning methods operate, thereby laying the groundwork for our NN–based cleaning procedure.

Let \widehat{C}_{MLE} be the MLE of the correlation matrix obtained from Δt_{in} observations drawn from a multivariate distribution with covariance matrix $\Sigma = DCD$, with $D = \text{diag}(\sigma_1, \dots, \sigma_n)$ being the diagonal matrix of the population assets' volatilities and C the population correlation matrix, and let $\widehat{C}_{\text{MLE}} = \widehat{V} \widehat{\Lambda} \widehat{V}^{\top}$ be the spectral decomposition of the MLE. A RIE replaces the noisy eigenvalues by a deterministic function f while keeping \widehat{V} unchanged:

$$\widehat{\Xi}_f := \widehat{\mathbf{V}} \, f(\widehat{C}_{\mathsf{MLE}} \mid q) \, \widehat{\mathbf{V}}^\top. \tag{5}$$

Assuming rotational invariance of the sampling process, the function f can be expressed in terms of the eigenvalues of the empirical correlation matrix

$$f(\widehat{C}_{MLE} \mid q) \longrightarrow \operatorname{diag}(f(\widehat{\lambda} \mid q)).$$
 (6)

Notably, f is a vector-valued function, i.e., $f: \mathbb{R}^{n+1} \longrightarrow \mathbb{R}^n$, such that

$$f(\widehat{\lambda}_1, \dots, \widehat{\lambda}_n \mid q) = \left\{ f_1(\widehat{\lambda}_1, \dots, \widehat{\lambda}_n \mid q), \dots, f_n(\widehat{\lambda}_1, \dots, \widehat{\lambda}_n \mid q) \right\}, \quad \text{with} \quad f_i : \mathbb{R}^{n+1} \to \mathbb{R};$$
 (7)

which means that it is generally a non-separable map since each component may depend on the entire argument. Furtermore, f is permuational-equivariant; therefore, called $\tau \in S_n$ a permuation operator, then

$$\tau\left(f(\widehat{\lambda}|q)\right) = f\left(\tau(\widehat{\lambda})|q\right), \quad \forall \tau \in \mathcal{S}_n.$$
 (8)

These two structural properties of f (Eqs. (7)–(8)) anticipate the constraints we impose on the NN architecture in Sec. 3.2.

The optimal $f^*(\hat{\lambda} \mid q)$ is defined as the minimiser of the Frobenius-norm loss against the (unknown) population correlation matrix,

$$f^{\star}(\widehat{\lambda} \mid q) := \arg \min_{\lambda \in \mathbb{R}^n} \left\| C - \widehat{V} \operatorname{diag}(\lambda) \widehat{V}^{\top} \right\|_F^2.$$
 (9)

Because it requires knowledge of the (unknown) population correlation matrix C, the exact solution of Eq. 9 is called an oracle $f^{OE}(\hat{C}, C)$ and is not implementable in practice.

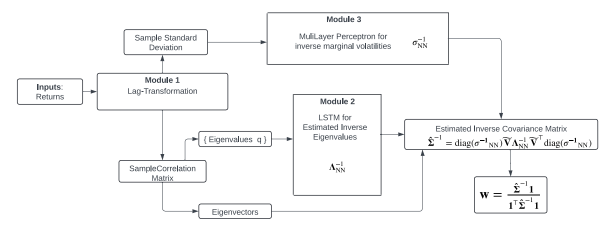


Figure 1: Schematic representation of the proposed NN architecture.

A straightforward minimisation of Eq. (9) yields the closed-form solution

$$f_i^{\text{OE}}(\widehat{\boldsymbol{C}}, \boldsymbol{C}) = \sum_{j=1}^n \lambda_j \, \psi_{ij}, \qquad \psi_{ij} := \sum_{k=1}^n (\widehat{v}_{ki} v_{kj})^2.$$

$$(10)$$

where ψ_{ij} is the overlap between the *i*-th eigenvector of the empirical correlation matrix and the *j*-th eigenvector of the population correlation matrix. Given that Ψ is an overlap between two reference frames, it is trivial to prove that it is a bi-stochastic matrix, i.e., the sum of each row and each column is equal to one. From this property comes that the oracle eigenvalues can be viewed as a weighted average of the population eigenvalues, where the weights are given by the overlap between the empirical and population eigenvectors. For bulk indices *i*, the weights ψ_{ij} are widely dispersed over the *j* elements in the bulk, so that f_i^{OE} becomes an almost uniform average of many population eigenvalues within the bulk. In a rough approximation, eigenvalue cleaning replaces the noisy fine structure of the bulk by a single effective level, while leaving the well-isolated high eigenvalues nearly untouched.

Clearly, this estimator is not available in practice because C is unknown. However, under rotational invariance, the expectation of both the population eigenvalues Λ and the overlap Ψ must depend only on the eigenvalues of the empirical covariance matrix $\hat{\Lambda}$ and the aspect ratio $q=n/\Delta t$. Such expectation cannot be derived analytically in a closed form, but NLS theory provides an analytical approximation of $f^* \to f^{\rm OE}$ in the limit $n, \Delta t \to \infty$ with q fixed [13].

It is critical to notice that the loss function (Eq. (9)) used in the NLS theory is the Frobenius distance between the population correlation matrix and the empirical correlation matrix. This is not the same as the loss function used in a GMV portfolio optimization, which should be the realized OOS variance of the portfolio weights σ_{PTF}^2 . Although in Ref. [45] the authors claim that both loss functions are equivalent, this is not true in practice. In Ref. [33], it was shown that the Frobenius-based OE is, in fact, not optimal for portfolio optimization.

3 Neural Networks for Portfolio Optimization

The NN architecture that we propose is organised to replicate the analytical workflow of the GMV estimator. Embedding the algebraic symmetries of the GMV solution into the network topology both limits the number of free parameters and promotes robust OOS performance in large universes.

Fig. 1 highlights the three learnable modules of our architecture. First, the lag-transformation module (Model 1, Sec. 3.1), applies a learnable temporal weighting kernel and a soft-clip to the raw return matrix R, yielding the transformed returns \widetilde{R} . Different lags can receive unequal weights; recent observations may be stressed, older observations may be down-weighted, or the reverse, depending on what minimises the loss. The resulting sequence bifurcates into two parallel branches.

The upper branch standardises $\widetilde{\boldsymbol{R}}$ to $\widetilde{\boldsymbol{Z}}$ and passes it to the eigenvalue-cleaning module (Model 2, Sec. 3.2). This block learns a non-linear shrinkage of the sample correlation matrix $\widetilde{\boldsymbol{C}} = \widetilde{\boldsymbol{Z}} \widetilde{\boldsymbol{Z}}^\top / \Delta t_{\rm in}$, producing a cleaned inverse eigenvalue matrix $\boldsymbol{\Lambda}_{\rm NN}^{-1}$.

The lower branch processes the vector of marginal volatilities $\{\widetilde{\sigma}_1,\ldots,\widetilde{\sigma}_n\}$ of \widetilde{R} through the marginal-volatility network (Model 3, Sec. 3.3). This network transforms each standard deviation into its optimal inverse, generating the diagonal matrix $D_{\mathrm{NN}}^{-1} = \mathrm{diag}(\sigma_{\mathrm{NN}}^{-1})$.

The two branches recombine to form the estimator of the inverse covariance matrix,

$$\Sigma_{NN}^{-1} = D_{NN}^{-1} \tilde{V} \Lambda_{NN}^{-1} \tilde{V}^{\top} D_{NN}^{-1}, \tag{11}$$

where \widetilde{V} contains the eigenvectors of \widetilde{C} . Substituting Σ_{NN}^{-1} into $\widehat{\Sigma}^{-1}$ of Eq. (1) yields the portfolio weights.

The architecture is trained end-to-end by minimising the realised OOS variance of the resulting portfolio. The loss function is

$$\mathcal{L}(\boldsymbol{w}, \boldsymbol{\Sigma}_{\text{out}}) = n \, \boldsymbol{w}^{\top} \boldsymbol{\Sigma}_{\text{out}} \boldsymbol{w}, \tag{12}$$

where Σ_{out} denotes the OOS realized covariance matrix of portfolio returns. The factor n rescales the objective so that its magnitude is less sensitive to the cross-sectional dimension.

3.1 Lag-Transformation module

It is reasonable to assume that past observations do not influence the future risk uniformly. To make our NN lag aware, we design a specific module that, before the input returns reach the subsequent blocks, they are processed through a learnable lag-transformation system that assigns a distinct transformation to every look-back lag. Classical risk models define a fixed exponential kernel [46, 20]; our formulation generalises that kernel by letting the NN determine its shape by training on the real data.

Each raw return r_{ti} is mapped to

$$\widetilde{r}_{ti} = \frac{\alpha_t}{\beta_t} \tanh(252 \,\beta_t r_{ti}),\tag{13}$$

with $\alpha = \{\alpha_t\}_{t=1}^{\Delta t_{\rm in}}$ and $\beta = \{\beta_t\}_{t=1}^{\Delta t_{\rm in}}$ learned by back-propagation. The coefficient α_t rescales observations at lag t. The coefficient β_t modulates their outlier sensitivity by controlling the saturation of the hyperbolic tangent. When β_t is small the mapping is nearly linear $(\tanh(x) \approx x \text{ if } x \text{ is small})$ and the return enters almost unchanged; when β_t is large the mapping approaches a symmetric step that clips extreme values at $\pm \alpha_t/\beta_t$. In this way, the module can emphasise or attenuate individual lags and dampen outliers simultaneously.

The factor 252 annualises the input and stabilises gradient computations under single-precision arithmetic. Because α_t and β_t are updated jointly with all other network parameters, the module can learn any monotone or non-monotone lag profile, with the classical exponential decay arising as a special case. The output transformed matrix $\widetilde{\boldsymbol{R}}$ serves as the input to all subsequent modules.

3.2 Eigenvalue Cleaning with Neural Networks

The second module replaces the Ledoit and Péché RIE with a purely data-driven alternative. It learns a vector-valued function $f: \mathbb{R}^{n+1} \to \mathbb{R}^n$ that transforms the empirical eigenvalues $\{\widetilde{\lambda}_1, \dots, \widetilde{\lambda}_n\}$ of the lag-transformed returns' correlation matrix (Eq. (13)) into the cleaned spectrum $\{\lambda_1^*, \dots, \lambda_n^*\}$.

As shown in Eqs. (7)-(8), the target function f is both non-separable and invariant under any permutation of its inputs. Applying a fully connected layer directly to the unordered vector of empirical eigenvalues would require handling O(n!) possible input orderings, which is plainly infeasible. Deep Set architectures [47] and attention-based models [48] enforce permutation symmetry and thus avoid this factorial blow-up, yet neither fully exploits the structure of the problem. Deep Sets adopt a top-down view: each eigenvalue interacts with a global pooled representation of the spectrum. Although universal approximation theorems guarantee expressiveness [49], modelling fine interactions would require an exploding number of pooling tokens as n increases. Attention mechanisms implement a bottom-up approach that begins by computing pairwise interactions. Capturing higher-order dependencies then requires stacking multiple attention-feed-forward modules, which deepens the network, exacerbates vanishing or exploding gradients [50, 51], and poses significant computational challenges as n grows.

To design an architecture that more effectively mitigates the curse of dimensionality, we exploit the exact mathematical mapping provided by the Coulomb-gas formulation of RMT [34], which models sample eigenvalues as one-dimensional charges under three forces: mutual logarithmic repulsion, confinement by the population spectrum, and hard-edge repulsion at zero. The confinement potential reflects the full population spectrum: when eigenvalues are well separated relative to the effective sample length $\Delta t_{\rm in}$, it is dominated by the nearest population eigenvalue; otherwise, it arises from a complex superposition of bulk contributions. The mutual repulsion between sample eigenvalues scales as

 $|\widetilde{\lambda}_i-\widetilde{\lambda}_j|^{-1}$ and therefore decays with rank separation, while the hard-edge term enforces nonnegativity but weakens as $\Delta t_{\rm in}$ approaches n. Consequently, the shift of each $\widetilde{\lambda}_i$ away from its population counterpart is governed primarily by interactions with the sample eigenvalues that lie within its local spectral vicinity, a region whose extent depends on the local spacing around $\widetilde{\lambda}_i$, while the cumulative influence of all more distant eigenvalues can be approximated as a coarse-grained contribution. Although the resulting spectral density and eigenvector-overlap dynamics could in principle be derived by solving the coupled Hamiltonian equations associated with this analogy, we instead incorporate only the implied locality as an inspiration for a structural prior in our neural architecture, which should lie, in fact, between the extremes of Deep Sets and attention architectures.

To clarify the last statement, consider that within any subset of sample eigenvalues, the effective order of interaction is not fixed in advance but grows with the local density of the population spectrum. Regions with tightly spaced population eigenvalues induce strong collective forces that demand modelling higher-order dependencies among neighbouring sample eigenvalues, whereas sparse regions admit accurate approximation with only low-order interactions. In a bottom-up framework such as self-attention, capturing these varying orders requires explicitly stacking as many attention–feed-forward layers as the maximum local interaction order, since each layer only learns pairwise or limited higher-order dependencies. As the dimension n increases, a region of given spectral concentration spans more eigenvalues, forcing deeper networks to maintain the same modelling fidelity. Consequently, this scaling renders pure attention architectures impractical in very high dimensions.

A Recurrent Neural Network (RNN) can naturally capture such locality. Viewing the rank index as a pseudo-time coordinate, we process the ordered sequence $\widetilde{\lambda}_1 \leq \cdots \leq \widetilde{\lambda}_n$ with a Bidirectional Long Short-Term Memory (BiLSTM) network [52, 53], which is a specific architecture of the RNN family. An RNN employs a single learnable cell function, so that at each step it takes the current input and the previous hidden state to produce an updated hidden state. This same cell is applied repeatedly across the ordered sequence, allowing the model to process inputs of arbitrary length with a fixed set of parameters.

For the bidirectional version, the forward and backward cells learn the update functions

$$g^{\rightarrow}(\boldsymbol{h}_{i-1}^{\rightarrow}, \boldsymbol{m}_{i-1}^{\rightarrow}, \widetilde{\lambda}_{i}, q) = \{\boldsymbol{h}_{i}^{\rightarrow}, \boldsymbol{m}_{i}^{\rightarrow}\}, \quad \text{with} \quad i = 1, \dots, n,$$
 (14)

$$g^{\leftarrow}(\boldsymbol{h}_{i+1}^{\leftarrow}, \boldsymbol{m}_{i+1}^{\leftarrow}, \widetilde{\lambda}_i, q) = \{\boldsymbol{h}_i^{\leftarrow}, \boldsymbol{m}_i^{\leftarrow}\},$$
 with $i = n, \dots, 1,$ (15)

where $\boldsymbol{h}_{i}^{\rightleftharpoons} = \left\{h_{i,1}^{\rightleftharpoons}, \ldots, h_{i,k}^{\rightleftharpoons}\right\}$ and $\boldsymbol{m}_{i}^{\rightleftharpoons} = \left\{m_{i,1}^{\rightleftharpoons}, \ldots, m_{i,k}^{\rightleftharpoons}\right\}$ are the hidden and cell states. Setting the input states to zero at the start of the sequences, Eq. (14) is applied from the smallest to the largest eigenvalue, whereas Eq. (15) proceeds in the opposite direction.

The forward states summarise all eigenvalues to the left of the current rank, and the backward states summarise those to its right. Their concatenation, therefore, encodes a two-sided interaction whose effective range is governed by LSTM gating rather than by an explicit window. Using rank as the temporal axis is unconventional but consistent with the symmetry of long-range yet decaying interactions. By adopting a BiLSTM, the model no longer requires capacity that grows with n, as was necessary for both Deep Sets and attention mechanisms. Instead, the BiLSTM learns to approximate the combination of local neighbor interactions and a distant mean-field term. RMT indicates that this combined interaction converges to a fixed filter as n and $\Delta t_{\rm in}$ increase [14], so model complexity should remain bounded even in very high dimensions.

After scanning the spectrum, the BiLSTM outputs an $n \times 2k$ matrix H whose i-th row h_i concatenates column-wise the forward h_i^{\rightarrow} and backward h_i^{\leftarrow} hidden vectors. A dense layer shared across ranks projects h_i onto the positive real line through a softplus activation

$$\lambda_{i,\text{NN}}^{-1} = \text{softplus}(\mathbf{w}^{\top} \mathbf{h}_i + b), \qquad i = 1, \dots, n,$$
(16)

with trainable parameters $\mathbf{w} \in \mathbb{R}^{2k}$ and $b \in \mathbb{R}$. The softplus retains differentiability at the origin and avoids hard saturation, ensuring stable gradients even for very small eigenvalues. To remove the arbitrary overall scale, we normalise such that $\sum_{i=1}^n \lambda_{i,\mathrm{NN}}^{-1} = n$. Portfolio weights computed from a RIE do not change if all the eigenvalues are multiplied by the same constant, so this normalization places every instance on the same scale and allows direct comparison in downstream statistical tests without altering the resulting portfolio weights.

In summary, the module comprises a total of $8k^2 + 26k + 1 = 34,433$ learnable parameters (k = 64). Both LSTM cell contributes $8k^2 + 24k$ parameters, and the final dense layer adds 2k + 1. If model compactness is a priority, the LSTM units may be replaced by two Gated Recurrent Units (GRUs). Standard LSTMs employ separate memory and hidden channels, m and h, respectively, to mitigate vanishing and exploding gradients. The cell state m, modulated by input, forget, and output gates, can preserve long-range information almost unchanged over many time steps, while the hidden state h undergoes more aggressive step-wise updates and serves as the immediate output. This arrangement

improves numerical stability and captures both global and local spectral structure. That said, this dual-channel design is not strictly necessary: in our experiments, a simplified GRU, which merges long and short-term memory into a single h performed comparably well. The GRU reorganizes its gating mechanism so that the hidden state alone can reliably encode long-range dependencies, offering a lighter yet effective alternative. Substituting LSTM cells with GRUs reduces the total parameter count to $6k^2 + 19k + 1$.

3.3 Multilayer Perceptron for Inverse Marginal Volatilities

The second branch of the network estimates the inverse marginal volatilities from the sample standard deviation of the transformed return series. i.e.,

$$\widetilde{\sigma}_{i} = \sqrt{\frac{1}{\Delta t_{\text{in}}} \sum_{t=1}^{\Delta t_{\text{in}}} \widetilde{r}_{t,i}^{2} - \left(\frac{1}{\Delta t_{\text{in}}} \sum_{t=1}^{\Delta t_{\text{in}}} \widetilde{r}_{t,i}\right)^{2}}.$$
(17)

Each standard deviation is then passed through an MLP network with 3 hidden layers of (64, 32, 16) neurons and a Leaky-ReLU activation function. The output of the network is a single neuron with a softplus activation function, which guarantees that the output is positive. In total, this module comprises 2,753 learnable parameters. The output of this branch is then standardized by dividing it by the average inverse output of the network, i.e.,

$$\sigma_{i,\text{NN}}^{-1} \to \frac{\sigma_{i,\text{NN}}^{-1}}{\frac{1}{n} \sum_{j=1}^{n} \sigma_{j,\text{NN}}^{-1}}.$$
 (18)

This normalization is irrelevant for the portfolio weights, but it is useful to compare the output of the network across different samples.

Although a more sophisticated design could make the volatility transformation context-dependent, adapting to correlation structure or different market regimes, we deliberately adopt this minimal architecture. Our focus in this work is on the eigenvalue-cleaning of the correlation matrix, and hence, we keep the marginal volatility branch as simple as possible.

4 Training and Evaluation

4.1 Selection of the Investable Universe

We describe here the procedure used to select assets for investment from the universe of US equities that were listed on the NYSE or NASDAQ at any point from 1 January 1990 to 31 December 2024. This universe, therefore, includes American Depositary Receipts (ADRs) and explicitly excludes funds and exchange-traded funds. To avoid data leakage, for each trading day t, we rely only on information available up to t-1 when determining the eligible stocks. We further exclude any security for which a planned delisting occurs within the subsequent $\Delta t_{\rm out} = 5$ trading days.

We designed a two-tiered filtering scheme that operates on both the full five-year calibration window ($\Delta t_{\rm in}=1200$) and a short, recent time window. The full-history filters enforce homogeneity of trading activity over the entire five years and thus attempt to remove any stock whose price series is fragmented by late initial public offerings, prolonged trading halts, or significant corporate restructurings. The recent-window filters, by contrast, apply stringent liquidity thresholds over the last five to twenty trading days with the attempt to eliminate any security whose executed auctions fall below the levels compatible with low-impact rebalancing. This dual approach both addresses the challenges of assembling a broad universe of n=1000 stocks with uninterrupted five-year histories and ensures that each selected asset remains sufficiently liquid at the time of investment. That said, the threshold values themselves are inherently arbitrary and have been chosen primarily to maintain a stable universe of n=1000 stocks at every point in time; nevertheless, filtering approaches like this reflect standard practice in hedge funds, where inclusion criteria must satisfy stringent and complex rules. In practice, working with a very large, unconstrained dataset can introduce unintended biases and degrade the reliability of both risk estimates and optimized allocations.

Our full history filter enforces a homogeneous trading record across the five-year calibration window by requiring that, within any rolling one-year window, a stock has participated in the closing auction on at least 95% of trading days. By doing so, we guarantee that at every point in time within the year-long window, the vast majority of price observations are present.

On the recent trading history, we require that 100% of the auctions are executed and at least 1% of the shares outstanding and 1% of the market capitalisation are traded on average over the last five days. This ensures that the stock is liquid enough to be traded with a low impact on the market. The day before the trade, we check that the number of shares outstanding is greater than 5M, to avoid micro-shares. Finally, on the day before trading, we require that each stock's

price lies between 10 \$ and 2,000 \$. Stocks trading at relatively low price levels incur high per-share transaction costs, whereas those at relatively high price levels often exhibit reduced trading activity, impeding efficient execution.

Since our aim is to design a multivariate methodology to obtain a portfolio of stocks with a minimum variance, we decided to filter out those stocks that univariately exhibit an extremely low variance when compared to the universe of stocks. In the presence of those cases, the minimal variance becomes trivial, and the multivariate nature of the problem is lost. In other words, we are interested in building a portfolio of high-variance stocks, which, when combined, can lead to a low-variance portfolio. To this end, we applied a standard procedure to remove outliers. We removed from the available universe those stocks whose distribution of log standard deviation is below the 1.5 interquartile range (IQR) of the log standard deviation of the universe [54]. This condition must be satisfied in two time windows of 5 and 20 most recent days.

Finally, to reduce redundancy, we keep only one class of shares for each company, the one with the largest market capitalization on the previous day. A similar filter is executed for those stocks with an in-sample correlation greater than 0.95 [55]. The final selection is done by ranking the stocks by their market capitalization and selecting the top n=1,000 stocks for every trading day t. The final panel, spanning January 1990 to December 2024, contains roughly 2.7 millions of stock–day pairs and 3, 837 unique stocks.

4.2 Training Procedure

The proposed architecture is designed to be independent of the number of assets n. This means that the same model can be applied to different subsets of assets without the need for retraining. However, we know from RMT that the filtering should depend on the aspect ratio $q=n/\Delta t_{\rm in}$; therefore, during the training phase, the NN should be fed with examples of different n if we aim to obtain a model that generalizes well to larger portfolio sizes. This is done by randomly and uniformly sampling the number of assets n from a [50,350] range on every batch of the training. The number of days $\Delta t_{\rm in}$ is fixed to 1200 trading days, in fact, the presence of the module of Sec. 3.1 includes $2\Delta t_{\rm in}$ trainable variables (α_t,β_t) , which are associated to specific past lags. In the future version, this could be generalized to a variable number of past lags, but it would require a more complex architecture.

We trained our model on a dataset of US equities spanning January 1990 to December 2024. For each validation date t, we define the calibration window as the period from the dataset start through December 31 of the year preceding t. In particular, we conduct validation over the interval [2000-01-01, 2024-12-31], resulting in 24 distinct trained NNs. To construct a robust training procedure, we randomly draw a date t such that the interval $[t - \Delta t_{\rm in} - 1, t + \Delta t_{\rm out}]$ is fully contained within the calibration window. We then randomly select n assets from the investable universe at t-1 (see Section 4.1). The NN is trained on the adjusted close-to-close returns over the input window $[t - \Delta t_{\rm in}, t - 1]$ together with the realized OOS covariance matrix $\Sigma_{\rm out}$, computed from the adjusted close-to-close returns over $[t, t + \Delta t_{\rm out}]$. A one-day shift between the training inputs and the OOS returns is introduced to avoid data leakage, since the return at t depends on the closing price at the end of day t, which is not available intra-day.

According to the training procedure, the probability that NN will be trained on the same dataset is very low for n sufficiently smaller than the investable universe on a given day. This allows us to train the NN without the need to monitor the performance on a validation set for early stopping. However, to further reduce the risk of overfitting, we used a decay factor of the learning rate, which is set to 10^{-4} and drops every batch by a factor of $0.99^{1/500}$, and we clip the gradients by norm to 1.0. The gradient is then optimized with the Adam optimizer with an automatic rescaling of the loss function. We trained all our models with 100 epochs, with 500 steps per epoch, and a batch size of 32 samples. To ensure that the convergence is stable, for each of the 24 years, we trained our NN 10 times independently, so finally, we obtained 240 trained NNs.

For further evidence of robustness against overfitting, Fig. 2 displays the evolution of training and validation losses over successive epochs. The loss trajectories remain consistent across independent runs on the same data interval. Both training and validation losses decrease monotonically. Monotonic decline of the training loss is expected; the validation loss, by contrast, often increases once overfitting sets in. After 60 epochs, the training loss plateaus, exhibiting an average change of less than 3.5% over the final forty epochs. The validation loss continues to decline at a faster pace, falling by approximately 6% during the same period. These results indicate that terminating training around 100 epochs would probably incur minimal overfitting risk.

4.3 Model Interpretability

4.3.1 Lag-Transformed Returns

After the calibration, we can analyze the learned weighting factors α_t and β_t of Eq. (13) as a function of the lag t to understand how the NN is weighting and transforming the past returns. Given that we ran 10 independent training runs

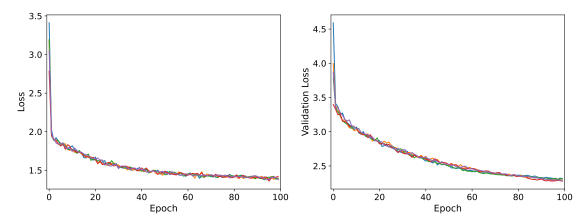


Figure 2: Training loss on the left panel, validation loss on the right panel. Different lines refer to independent training runs on the same calibration window. The figure refers to the calibration of the first model up to 31-12-1999. The loss function is defined in Eq. (12).

for each of the 24 years of the validations, we obtained 240 trained models. We can then compute the average of the weighting factors over the trained models, which are shown in Fig. 3.

Interestingly, the rescaling factor α_t seems to follow a power law with an exponent of approximately -0.2, which in substance means that the most recent past is weighted slightly more than the oldest past. In fact, half of the contribution is given from the 40% most recent past. The emergence of the power law is quite surprising, being the most widely used decay factor based on Exponentially Weighted Moving Average (EWMA) [46, 20], whereas the power law decay would instead imply a Hyperbolic Weighted Moving Average (HWMA). A model based on HWMA was proposed in Ref. [56], but to our knowledge, it is not very popular yet.

The clipping factor β_t shows values within the range [0.4, 1.0]. Interestingly, the clipping factor β_t seems to saturate for larger lags. To have an idea about the range of clipping, for the farthest data points, assuming $\alpha_t = 1$, a daily return of 1% is clipped to approximately 0.4% while the same return in the most recent past is clipped to approximately 0.9%.

The lower panels of Fig. 3 present the combined impact of the shrinkage and sigmoid transformations. In the remote past, the transformed values not only contract toward zero but also assume effectively binary states. As a result, beyond approximately one year of lag the series conveys only the sign of daily returns. When the Pearson correlation is applied to a binarized series, it coincides with the Phi coefficient [57, 58]. In contrast, the most recent returns still exhibit a strong binary character yet develop a pronounced uniform region around the center. This feature reflects the fact that the sigmoid kernel closely tracks the empirical cumulative distribution. By transforming the raw data via the empirical CDF, the Pearson correlation of the transformed series becomes the Spearman coefficient [59]. However, this exact equivalence holds only when each return series is rank-transformed by its own empirical CDF, whereas using a single merged CDF across all assets only approximates the Spearman coefficient and remains inexact due to volatility-driven distortions across assets. In summary, as the lag increases, the network's operative correlation metric shifts continuously from a Spearman-type measure at short lags toward a Phi-type measure at long lags.

4.3.2 Eigenvalue Sensitivity Analysis

The surprising result of the AO estimator [30, 29] is that a set of fixed eigenvalues can be used to obtain an estimator for the correlation matrix that outperforms all the RMT-based estimators for portfolio optimization. Moreover, such an estimator does not depend on the assets used in the calibration or the exchange, country, or time period. The authors of Ref. [30] hypothesized that the non-stationarity of the system is so strong that, unless eigenvectors are corrected, very little information can be extracted from the sample eigenvalues. In our case, we used an NN trained to attempt extracting specifically the non-stationary relationship between $\hat{C}_{\rm in}$ and $C_{\rm out}$. If the above statement is true, and the NN does not overfit the calibration data, we should expect an output that is independent of the assets in the input.

In the left panel of Fig. 4, we present the eigenvalue spectra for the MLE, the QIS shrinkage, and the NN-transformed eigenvalues λ_{NN} after inversion of Eq. (16). The range of the largest eigenvalues coincides across all three estimators, whereas the bulk of the spectrum exhibits the most pronounced differences. The MLE assigns the lowest values to the bulk eigenvalues. The QIS shrinkage attenuates this dispersion by smoothing the spectrum. Neither method reproduces the abrupt truncation characteristic of eigenvalue clipping [9]; instead, while both the MLE and QIS yield a bulk

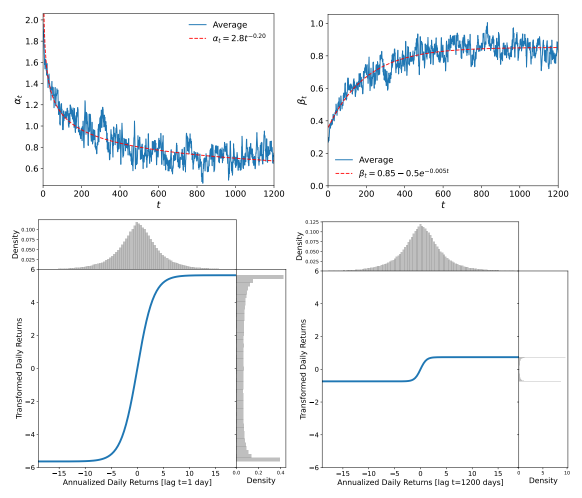


Figure 3: The upper plots show the calibrated weighting factors α_t and β_t as a function of the lag. The factors shown here are an average over the 240 trained models. Low values of the x-axis correspond to the most recent past, while high values correspond to the oldest past. Both figures show an attempt to fit them with a power law for α_t and with a saturating exponential for β_t indicated with the dashed red lines. The lower plots show an example of the return transformation of the annualized returns for the last trained model over past examples. The left plot refers to the day before the investment, and the right plot refers to the first day in the dataset. The marginal distribution shows how the empirical distribution is transformed.

spectrum spanning approximately one order of magnitude, the NN transformation compresses the bulk eigenvalues to within roughly half an order of magnitude. This compression causes the NN transformation to resemble more the abrupt truncation of eigenvalue clipping.

The right panel of Fig. 4 shows the standard deviation of the log-transformed eigenvalues across calibration samples. All estimators exhibit pronounced variability for the largest eigenvalues. In the bulk of the spectrum, the MLE and the QIS continue to display substantial dispersion, whereas the NN-transformed eigenvalues form a nearly flat plateau. This plateau indicates that the NN mapping renders the bulk eigenvalues effectively insensitive to the input data, akin to the asset-agnostic behavior of the AO estimator, while still responding to the extreme eigenvalues.

4.3.3 Marginal Volatility Estimator

The second branch of the NN architecture is a deep NN that learns the inverse marginal volatilities. After the training, we can analyze how the NN transforms the standard deviation of the rescaled returns. It is important to notice that this transformation is applied to every standard deviation independently of other stocks and market conditions, and it is also not dependent on the history of the stock itself. Although we believe that this is not the optimal way to learn this variable, the main task of this work is the learning of the eigenvalue cleaning, and this module is set up by purpose as

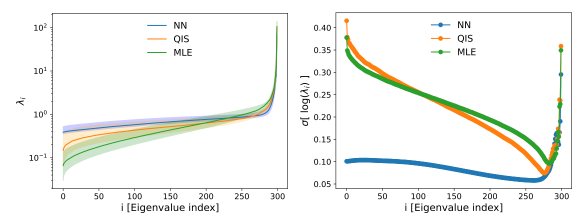


Figure 4: Eigenvalue sensitivity analysis. Left: median of the eigenvalues as a function of the rank, the colored bands indicate a 95% percentile coverage. Right: Standard deviation of the log eigenvalues as a function of the rank. The eigenvalues are computed on 1024 samples from the full calibration dataset.

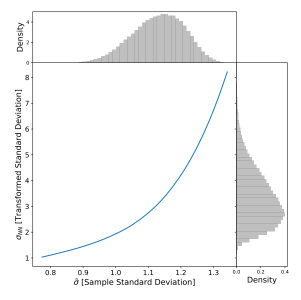


Figure 5: The figure shows how the Deep NN (Model 3) transforms the standard deviation of the rescaled returns. The x-axis shows the standard deviation of the rescaled returns, while the y-axis shows the output of the NN. The marginal distributions of the input and output are shown on the top and right sides of the figure, respectively.

minimally as possible (likewise, the returns lag-transformation module). In Fig. 5, we show how the NN transforms the standard deviation of the rescaled returns. The x-axis shows the standard deviation in input, and the y-axis shows the output of the NN. The marginal distributions of the input and output are shown on the top and right sides of the figure, respectively. Practically, the NN flattens the left tail of the distribution, which corresponds to low volatility assets, while it amplifies the right tail of the distribution, which corresponds to high volatility assets. This means medium-low volatility assets will have a marginal decrease in volatility, while high volatility assets will have a marginal increase in volatility.

4.4 Backtesting Procedure

The calibrated network is trained to predict the GMV portfolio by allowing unrestricted short positions. This design choice yields a closed-form mapping of the optimal weights via the analytical expression in equation Eq. (1). If one imposes a non-negativity constraint on the weights, the GMV solution must be obtained through a numerical solver. Although specialized NN layers can implement such constrained quadratic programs with end-to-end differentiation [60], this approach incurs substantially higher computational cost and, in our tests, suffers from vanishing gradients. On

the other side, allowing short sales solely to minimize variance might be undesirable, since it amplifies gross leverage and incurs additional borrowing and transaction costs.

To reconcile analytical tractability with realistic trading constraints, after the uncontrained calibration, we can alter the architecture so that the network directly outputs an estimate of the inverse covariance matrix, Σ_{NN}^{-1} (see Fig. 1). We invert this matrix and then compute portfolio weights under a long-only constraint by solving the QP externally.

In Sec. 4.4.2 we evaluate our method against alternative estimators defined in Sec. 4.4.1 under idealized, frictionless conditions. In this experiment, we calibrate the network without trading constraints, i.e., allowing both long and short positions. Then, we test both the raw uncontrained output and the long-only version described above. In Sec. 4.4.3 investigates the model's extrapolation performance. We train on a small universe of assets and then test on a substantially larger universe, enforcing long-only positions and employing an operational simulator that incorporates realistic transaction costs.

4.4.1 Benchmarking Alternative Estimators

For clarity, we distinguish between methods that rely exclusively on univariate information and those that exploit the full covariance structure.

Univariate benchmarks serve as low-information baselines. The Inverse-Volatility Portfolio (IVP) assigns each asset a weight proportional to the reciprocal of its estimated variances, thereby equalising the contribution of each asset to total ex-ante volatility; the idea was popularised by Maillard, Roncalli and Teiletche [61]. The capitalisation-weighted portfolio (MCAP) coincides with the market proxy used in empirical asset-pricing work and therefore absorbs only the cross-sectional information embedded in free-float market values.

Multivariate benchmarks differ only in the way the $n \times n$ correlation matrix \hat{C} is obtained. After the correlation matrix is estimated, we always project on the univariate volatilities $\hat{\Sigma} = \hat{D}\hat{C}\hat{D}$ and we apply the QP optimization. Filtering the correlation rather than the covariance is a necessary step to put RIEs in their best performance setup.

The simplest method is the sample MLE uses the unfiltered sample correlation matrix computed over the look-back window. The Power Mapping (PM), introduced by Guhr and Kälber [62], suppresses estimation noise by raising every off-diagonal element of the de-meaned correlation matrix to a power $\gamma > 1$ and re-normalising; following their empirical recommendations, we set $\gamma = 1.2$. For the LS, we use the standard implementation of Sklearn [63] where the parameter is set up according to the original work of Ledoit and Wolf [10]. For QIS, we used a Python implementation released in the GitHub repository [64] that we adapted to improve the computational performance. Finally, AO is calibrated using the universe of the stocks as defined in Sec. 4.1 within 1990-2000. Then the eigenvalues are kept constant for the whole validation period, 2000-2024, independently of the stock in the basket.

With the exception of the MCAP, which is rebalanced monthly, all the other portfolios are rebalanced at the same weekly frequency as the NN strategy, and they are compared on the same set of stocks.

4.4.2 Frictionless Interpolation Backtesting

The first backtesting procedure we used aims to evaluate the performance of the NN within the range of the calibration setup. To avoid confusion, it is important to note that we still mean that the NN is trained in the past and tested in the future, as explained in Sec. 4.2. Instead, we mean that in our training, we explored a range of $n \in [50, 350]$ assets, and we are testing the NN on n = 300 assets that are within the range of the training set. This implies that the NN will interpolate what it has learned during the training phase, and it will not extrapolate to a larger number of assets. Moreover, in this backtesting procedure, we will not include any transaction costs, no slippage, no borrowing costs, or perfect fractional shares. This is the most coherent condition to evaluate if the trained NN was successful in learning the task. In the next section, instead, we will evaluate the NN on a larger number of assets, and we will include realistic conditions.

To assess in a statistically robust way the performances, we performed a battery of 1,000 independent backtests. Specifically, every simulation consists of approximately 5 years of data, starting from a random day t within [2000-01-01, 2019-12-31], and performing 250 portfolio rebalancing every 5 trading days. Our model will use the last available calibration obtained the year before the trading day so that no data leakage occurs. The list of assets is randomly sampled from the investable universe (see Sec. 4.1) on the first day of the simulation, and it is kept fixed if possible for the entire backtest. If some of the assets are excluded from the investable universe during the backtest, they are removed from the portfolio, and another random asset is added to the portfolio so that the number of assets remains constant.

In Tabs. 1 and 2 we report the average performances of the models tested on the same set of assets. The unconstrained (long–short) results in Tab. 1 show that the NN achieves the highest SR (1.01) with an annualized volatility of 10.9%

Method	SR	Volatity	Mean Return	$n_{ m eff}$	Turnover	Gross Leverage
NN	1.011	0.109	0.110	15.2	1.125	3.07
QIS	0.942	0.111	0.105	10.6	0.440	3.81
AO	0.907	0.121	0.110	28.6	0.163	2.13
PM	0.902	0.114	0.102	10.3	0.534	4.00
LS	0.893	0.115	0.103	7.2	0.708	4.74
MLE	0.868	0.117	0.102	6.5	0.785	5.01

Table 1: Unconstrained long-short portfolio backtests. The table shows the average performances of 1,000 independent simulations of the experiments with 300 stocks within [2000-01-01, 2019-12-31] performing 250 portfolio rebalancing every 5 trading days, without transaction costs. Values are sorted according to the Sharpe Ratio (SR), and bolds represent statistically significant top rank according to a percentile bootstrap with p-value smaller than 10^{-6} . Portfolio volatility, mean returns and SR are annualized, $n_{\rm eff}$ is computed by using the inverse Herfindahl index [65].

Method	SR	Volatility	Mean Return	$n_{ m eff}$	Turnover
NN	0.792	0.135	0.107	16.5	0.469
AO	0.740	0.146	0.108	33.2	0.098
LS	0.723	0.142	0.103	13.6	0.142
PM	0.721	0.143	0.103	17.6	0.131
MLE	0.721	0.142	0.102	13.3	0.143
QIS	0.719	0.142	0.102	13.5	0.138

Table 2: Constrained long-only portfolio backtests. The table shows the average performances of 1,000 independent simulations of the experiments with 300 stocks within [2000-01-01, 2019-12-31] performing 250 portfolio rebalancing every 5 trading days, without transaction costs. Values are sorted according to SR, and bolds represent statistically significant top rank according to a percentile bootstrap with p-value smaller than 10^{-6} . Portfolio volatility, mean returns and SR are annualized, $n_{\rm eff}$ is computed by using the inverse Herfindahl index [65].

and a mean return of 11.0%. The QIS estimator attains the second-best SR (0.94) but does so with a notably high gross leverage of 3.81, compared with the NN leverage of 3.07. The AO method ranks third in SR (0.91) yet operates with the lowest leverage (2.13) and minimal turnover (16.3%). These performance gains for the NN and QIS come at the cost of trading intensity. The network rebalances more than its entire notional every five trading days (turnover 112.5%), and QIS rebalances 44.0% of its portfolio at each rebalance. Such high turnover and leverage would incur significant financing, borrowing, and market-impact costs not captured in this frictionless experiment.

Once short-selling is forbidden (Tab. 2), most of the models perform similarly. This is a known behavior of covariance cleaning for portfolio optimization [26, 44]. The annualized volatility is around 14.2% for all the models, including the MLE covariance estimator, which is the non-cleaned covariance matrix. The AO estimator performs slightly worse, reaching 14.6% annualized volatility. The NN instead reaches 13.5% annualized volatility, which is the lowest among all the models by a substantial margin. In terms of the SR, the NN reaches a 0.79, which is the highest among all the models, while the AO estimator reaches the second-best SR of 0.74, while all the other methods range around 0.72. We want to stress that checking the SR is very important when measuring the performance of a portfolio in a large, uncontrolled setting. In fact, it might be possible that the data cleaning included stocks that must be excluded, for example, stocks that are about to be delisted or merged or stocks that are not liquid enough. In those cases the asset might have a very low volatility and a very low return. A portfolio that includes those assets will have a very low volatility, but it will also have a very low return, leading to a very low SR. The objective of a GMV portfolio is, in fact, to minimize the volatility of investing in risky assets.

The other two columns in Tab. 2 report the average number of effective assets in the portfolio. This metric is dominated by the AO estimator, which has an average of 33.2 effective assets, while the others range between 13.3 and 17.6, which are less than half of the number of assets in the portfolio.

Finally, the average turnover reported in the last column measures the fraction of assets that are changed in the portfolio at every rebalancing. The AO estimator has the lowest turnover (9%) while the NN has a turnover of 47%, which is the highest among all the models. This is due to the fact that the NN is trained to be reactive and to focus on the most recent past, while the other models are more stable and less sensitive to recent changes. This behavior is a double-edged sword; on one side, having a more reactive portfolio allows one to adapt to the market conditions, but on the other side,

it leads to a higher turnover and, therefore, higher transaction costs. For this reason, in the next section, we will evaluate the performance of the NN in a more realistic setting.

4.4.3 High Realistic Extrapolation Backtesting

The aim of this backtesting procedure is to evaluate the performance of the NN on a larger number of assets with long-only constraints and to include realistic conditions in our simulator. The main concern of the previous backtesting procedure (Sec. 4.4.2) is that the NN tends to have a high turnover, which can lead to high transaction costs and slippage.

The simulator described in this section reproduces, on a daily grid, the complete life cycle of an Interactive Brokers (IBKR) cash-and-margin account. All computations are performed in continuous calendar time, but market-sensitive events, price moves, dividends, corporate actions, and interest accruals are aligned to the exchange calendar supplied with the price data. The model is intentionally operational: every quantity that appears in a real IBKR statement (net liquidation value, margin excess, tiered credit and debit interest, commissions, SEC fees, clearing charges, etc.) is reproduced with the same algebra and the same day-count conventions used by the broker. No frictions are left unmodelled; conversely, no speculative optimization (such as VWAP execution, limit orders, or intraday crossing) is assumed so that the simulation yields a lower-bound estimate of attainable performance.

At any date t, the model keeps a position vector $\mathbf{s}_t = \{s_{t,i}\}_{i=1}^n$ expressed in shares and a free cash c_t scalar. Cash dividends are credited automatically and increase c_t . If free cash turns negative after trading, the account is deemed in debit. Interest accrues daily on the debit balance according to the IBKR tiered schedule [66]. Conversely, when c_t is positive, the broker pays credit interest on the portion above successive notional caps according to the corresponding credit-tier spreads; however, since the first 100,000\$ tranche is unpaid, we are not modeling that. To decide the daily rate a 360-day convention is used, so the daily debit rate is the FED funds rate $\phi_t^{(r)}$ [67] plus a spread δ_v , i.e., $\phi_t = \phi_t^{(r)} + \delta_v$. While the spread δ_v can be found on the IBKR website [68], the reference rate $\phi_t^{(r)}$ is a proprietary algorithm of the broker, and it is not available to the public. Nevertheless, we think that the FED funds rate could be a good proxy. It is worth noting that on the tiered schedule, the spread depends on the traded volume of the account in the current month.

At a rebalancing date, the target number of shares is determined from the estimate of Net Liquidation Value \widehat{NLV}_t . To compute the NLV, we need an estimate for the price of the equities at the closing auctions. We use the primary-exchange opening price if available; failing that, it defaults to the fully split-adjusted close of the preceding session. This choice mimics the information set of a manager who submits market orders before the end of the auctions, accepting execution price risk. The NLV pre-execution is then computed as

$$\widehat{\text{NLV}}_{t}' = c_{t-1} + \sum_{i=1}^{n} s_{t-1,i} \, \widehat{p}_{t,i}, \tag{19}$$

Please note that s_{t-1} is the position vector corrected by eventual splits. The target number of shares is then computed as

$$s_t = \text{round}\left(\frac{\mathbf{w}_t \, \widehat{\text{NLV}}_t'}{\widehat{\mathbf{p}}_t}\right).$$
 (20)

Finally, the actual NLV_t post-execution is computed with

$$NLV_t = c_t + \sum_{i=1}^{n} s_{t,i} \, p_{t,i}, \tag{21}$$

where p_t is the actual close price of the day, and c_t is already discounted for the dividend accrued during the day and fees.

The difference between the target number of shares s_t and the previous-day position s_{t-1} , corrected for eventual splits, is used as an end-of-day market order request $m_t = s_t - s_{t-1}$. The trades are executed with the actual close price of the day p_t , without accounting for market impact.

The difference between the target number of shares s_t and the previous-day position s_{t-1} , corrected for eventual splits, is used as an end-of-day market order request $\Delta s_t = s_t - s_{t-1}$. The trades are executed with the actual close price of the day p_t , without accounting for market impact. The executed volume is tracked to set up the proper tier of commissions and fees, and it is reset at the beginning of each month. Executed trades incur three additive costs. First, commissions follow the IBKR tiered schedule for US equities: a per-share rate of 0.035% when the month-to-date routed volume is below 300,000 shares and 0.020% above that threshold, with a floor of \$0.35 per ticket [66]. Second, exchange, clearing, and regulatory fees are modeled as a flat 0.0145% FINRA trading activity fee plus 0.05% exchange

	SR	Mean Return	Volatility	Max Drawdown	Turnover	$n_{ m eff}$
NN	1.046	0.122	0.117	-0.350	0.589	33.4
AO	0.951	0.118	0.124	-0.374	0.179	45.4
PM	0.880	0.109	0.124	-0.387	0.224	23.9
QIS	0.859	0.107	0.125	-0.388	0.230	19.0
MLE	0.858	0.107	0.125	-0.385	0.248	17.9
IVP	0.678	0.123	0.181	-0.504	0.058	728.0
MCAP	0.483	0.094	0.195	-0.542	0.065	133.1

Table 3: Performance of the long-only portfolio with n=1000 top capitalization stocks, rebalanced every 5 trading days, from January 2000 to December 2024. The simulations use a realistic trading simulator that accounts for transaction costs and slippage. SR, returns, and volatility are annualized. The turnover metric is computed without accounting for price drift and serves only as a reference to the model's stability, $n_{\rm eff}$ is computed by using the inverse Herfindahl index [65].

and 0.02% clearing fees, for a total of 0.0845% of notional executed. Third, "Section 31 SEC" fees apply to the sell notional at 1.157 basis points [66]. All charges are debited instantaneously from free cash.

The simulator returns, for every calendar date, the full path of NLV_t , cash, market values, and margin statistics, thus reflecting the realistic interaction between turnover, leverage due to slippage, and transaction costs. The simulation starts with a cash balance of 1,000,000 \$, which is a reasonable amount for a small hedge fund portfolio, and it is sufficiently low to avoid the need to account for market impact on the most liquid US stocks. Our portfolio is rebalanced every 5 trading days, and the investable asset universe is taken from the procedure described in Sec. 4.1, with the only difference that we select the top n = 1,000 for marketcap, defined from the previous day's close price. In this case, we are attempting to extrapolate the NN to a larger number of assets, given that the calibration was done on a maximum of 350 assets. Furthermore, $q = n/\Delta t_{\text{in}} \approx 0.83$, which is substantially larger than the training range of $q \in [0.04, 0.29]$.

In Tab. 3, we can observe the performance metrics of the portfolio across different strategies. The NN strategy outperforms the others in terms of SR, volatility, and maximum drawdown, and reaches the second-best mean return. The second-best SR is achieved by the AO estimator. All the other methods, including the MLE covariance estimator, have similar performances in terms of SR and volatility. The univariate methods, such as IVP and MCAP, perform significantly worse than the multivariate methods, with an SR of 0.678 and 0.483, respectively. In fact, for the IVP strategy, which allocates weights by inverting individual variances in a univariate fashion, the diversification indicator n_{eff} reaches 728 out of 1 000, reflecting an almost equally weighted allocation. As detailed in Sec. 4.1, stocks with exceptionally low IS volatility were excluded, thereby preserving the multivariate nature of the optimization problem.

In the top left panel of Fig. 6, we show the NLV of the portfolio as a function of time. The NLV is computed as in Eq. (21) and it is shown in log scale. From the figure, we can observe that the better performance of the NN in terms of returns is even higher after 2012. This is very important since many NN-based approaches tend to outperform only in the far past, typically before the huge surge of artificial intelligence [69], but we observe the exact opposite. In the plot, we just focus on four models, the NN, AO, PM, and MLE, which are the ones that are most different from each other according to the performance metrics in Tab. 3. In the top right panel of Fig. 6, we show the annualized volatility of the NN and AO portfolios as a function of time over a yearly rolling window. The NN portfolio has a systematically lower volatility than AO across almost the entire period, with the only exception of 2020, which is the year of the COVID-19 pandemic. In the bottom panel of Fig. 6, we show the Maximum Drawdown (MDD) of the portfolio by year. Also, this plot confirms that the NN portfolio has a systematically lower MDD than AO, with only a few exceptions.

5 Discussions

Recent studies discussed in this work illustrate the potential of neural networks in portfolio optimization. They capture nonlinear dynamics, enforce positive definiteness, combine convolutional and recurrent layers, or apply stagewise screening and weighting. They nevertheless share some limitations. They typically operate on low-dimensional or static universes that reduce generality. They employ heterogeneous benchmarks that often omit the GMV portfolio. They may leak future information through survivor-bias filters or delisting exclusions. Finally, the stability of these methods under varying cross-sectional dimensions remains under-investigated, leaving it unclear whether they truly maintain robust performances.

We aim to address these gaps by embedding each analytical step of the GMV solution into a modular neural architecture that remains agnostic to the number of assets. Our network learns the inverse covariance matrix end-to-end by

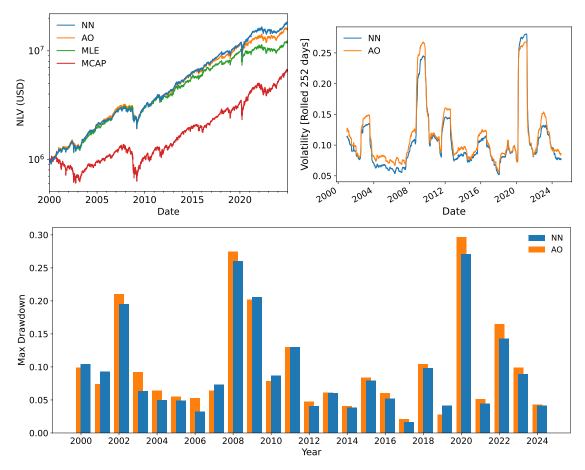


Figure 6: Long-only portfolio performances of the top 1000 most capitalized stocks in the universe backtested with the high realistic simulator. Top left: NLV of the portfolio as a function of time. Top right: Annualized volatility of the portfolio as a function of time over a yearly rolling window. Bottom: MDD of the portfolio by year.

minimizing future portfolio variance. It is calibrated on a few hundred equities and then applied without retraining to one thousand US stocks. The investable universe is selected each day using history-based and liquidity filters that prevent data leakage and exclude assets with anomalously low variance. In backtests with a high fidelity simulator from January 2000 to December 2024 under realistic trading conditions, our model delivers lower realized volatility, smaller maximum drawdowns, and higher Sharpe ratios than state-of-the-art methods. Notably, its end-to-end long-short calibration does not pose a serious limitation. In fact, when long-only constraints are imposed on its learned covariance matrix via an external optimizer, its performance advantage over competing estimators remains intact.

The interpretability of our approach follows from its three learnable modules. The lag-transformation block discovers a hyperbolic decay of past returns and applies soft clipping so that recent observations yield a Spearman-type correlation while older returns collapse to a binary sign. The eigenvalue-cleaning block employs a bidirectional LSTM to perform optimal eigenvalue cleaning. The outcome of this filtering is an approximately input-agnostic shrinkage of the eigenvalue bulk, which is less sensitive than Random Matrix Theory predictions, while eigenvalue outliers are dynamically adjusted by the network. The marginal-volatility block uses a simple feed-forward network that flattens the lower tail of the volatility distribution and inflates the upper tail. These modules reveal how temporal lags, risk modes, and individual volatilities shape the portfolio, thereby providing theoretical insight into the model's behavior. This interpretability also ensures transparency, aligning with industry standards that discourage opaque black-box solutions.

For future work, we see four areas for further improvement. First, one can replace the rotation-invariant eigenvalue filter with a context-dependent cleaning that adapts to evolving market regimes. Second, denoising can be extended to eigenvectors, a task that hierarchical clustering methods already accomplish successfully [25, 22, 23]. Third, the modules could be coupled so that lag-transformation, eigenvalue filtering, and volatility scaling interact dynamically rather than operating independently. Finally, we plan to generalize our framework to the full mean-variance problem by

integrating return estimation with covariance learning in a single end-to-end network, possibly including differentiable optimization layers to handle non-closed-form objectives. This extension would offer a more exhaustive solution to portfolio selection.

Author Contributions

C.B. conceived the study, designed the NN architecture, developed the code implementation, conducted the numerical simulations, and drafted the manuscript. E.M. contributed to the code development, carried out the exploratory simulations, data analysis, prepared the initial draft of the manuscript, and participated in the revision. R.N.M. reviewed the methodology and results, participated in discussions to refine the analysis, and contributed to the manuscript revisions.

Acknowledgments

This work was performed using HPC resources from the "Mésocentre" computing center of CentraleSupélec and École Normale Supérieure Paris-Saclay supported by CNRS and Région Île-de-France (http://mesocentre.centralesupelec.fr/). E.M. acknowledges a fellowship funded by PNRR for the PhD DOT1608375 in Sistemi complessi per le scienze fisiche, socio-economiche e della vita of Catania University.

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