Portfolio Selection with Probabilistic Utility, Bayesian Statistics, and Markov Chain Monte Carlo

Pietro Rossi§, Massimo Tavoni‡, Flavio Cocco‡ and Robert Marschinski¶

§ ENEA-HPCN, Via Martiri di Monte Sole 4, 40100 Bologna, Italy
‡ Prometeia S.r.l, Via Marconi 43, 40100 Bologna, Italy
¶ Institute for Physics, University of Potsdam, Potsdam, Germany

E-mail: pietro.rossi@prometeia.it, rossi@bologna.enea.it, massimo.tavoni@prometeia.it, flavio.cocco@prometeia.it, robert.marschinski@pik-potsdam.de

Abstract.
We propose a novel portfolio selection approach that manages to ease some of the problems that characterise standard expected utility maximisation. The optimal portfolio is no longer defined as the extremum of a suitably chosen utility function: the latter, instead, is reinterpreted as the logarithm of a probability distribution for optimal portfolios and the selected portfolio is defined as the expected value with respect to this distribution. A further theoretical aspect is the adoption of a Bayesian inference framework. We find that this approach has several attractive features, when comparing it to the standard maximisation of expected utility. We remove the over-pronounced sensitivity on external parameters that plague optimisation procedures and obtain a natural and self consistent way to account for uncertainty in knowledge and for personal views. We test the proposed method against traditional expected utility maximisation, using artificial data to simulate finite-sample behaviour, and find superior performance of our procedure. All numerical integrals are carried out by using Markov Chain Monte Carlo, where the chains are generated by an adapted version of Hybrid Monte Carlo. We present numerical results for a portfolio of eight assets using historical time series running from January 1988 to January 2002.
Introduction

Classical portfolio selection \[26\] by Maximisation of Expected Utility (MEU) suffers from well-documented drawbacks \[27\]: it often leads to extreme and hardly plausible portfolio weights, which additionally are very sensitive to changes in the expected returns. Moreover, it does not take into account differences in the level of uncertainty associated with the various input variables (estimation-errors), since its straightforward optimisation procedure imposes infinite faith on the estimated parameters. Historical data provides some information on future returns, but it is well known that simple-minded use of this information often leads to nonsense because estimation disturbance overwhelms the value of the information contained in the data. In fact, the positions of extrema of a function are often highly sensitive to irrelevant distribution details and it is thus quite simple to build examples (see following section) where a minimal parameter variation induces a very large shift in the extrema location.

The issue of uncertainty in expected returns and its implications for portfolio selection has been extensively analysed in the relevant literature: starting with the work of Bawa, Brown and Klein \[5\], many authors have since addressed the problem, often resorting to a Bayesian framework \[4, 6, 22, 25, 15, 27, 12\]. More recently, with a growing debate on asset return predictability (which will not be addressed here), the issue has re-gained the attention of the academia \[2, 3, 7, 24, 18\].

Nevertheless, parameters determined by observation of historical data are not the only source of trouble for portfolios based on function optimisation: all the expected utility maximisation procedures suffer from the presence of a scalar parameter related to the investor’s risk aversion, whose value cannot be set by the theory but still sensitively affects the resulting portfolio composition. Actually, due to a complete lack of scale for this risk-aversion parameter, it is usually adjusted ex post by hand, i.e. by merely observing where “the dynamics happen” and defining an ad hoc scale according to the simple prescription “increase the parameter if you want a more aggressive - meaning riskier - portfolio”. In some cases this might be an acceptable “degree of freedom”, allowing to customise portfolios, but when combined with the very parameter sensitive maximise-expected-utility-optimisation (MEU in the following), it turns out to produce highly unstable and inconsistent portfolios, meaning that a portfolio might change significantly for an apparently small shift in risk-aversion, and might even be less “aggressive” than a neighbouring portfolio with a lower risk-aversion. This will be discussed in more detail in Section 4. To our knowledge, this relation between optimisation procedure and risk-aversion parameter has not been investigated in previous studies.

The primary objective of this paper is to offer a common prescription for easing both of these pathologies. In order to eliminate the intrinsic optimisation instability caused by the over-sensitiveness towards external parameters, we suggest a different interpretation of the utility function. We consider the utility function to be the logarithm of the probability density for the portfolio to assume a given composition, and we define as optimal the expected value of the portfolio’s weights with respect to that probability.
As will be shown, this leads to an improved, more robust portfolio selection procedure, which allows us to incorporate the risk-aversion parameter in a stable and - even if not theory determined - at least self-consistent manner.

As for the issue of uncertainty in parameter determination, we adopt a fully Bayesian approach, in which parameters characterising the distribution of the data are described by distributions themselves. Additionally, the Bayesian approach offers a natural framework for the incorporation of subjective investor views into the portfolio selection procedure. Finally, through this method uncertainty is taken into account by stating explicitly the errors associated with the determination of the portfolio.

In what follows, we first introduce and discuss a theoretical framework in general terms. When coming to the specification of the posterior distribution and of the utility function, we resort to a multivariate Gaussian distribution framework, in line with common practice, deferring relaxation of this assumption to future research.

The final contribution of our paper concerns the numerical technology employed to perform all the relevant integrals. Most of these cannot be computed explicitly and therefore we will resort to a dynamical Monte Carlo integration or “Markov Chain Monte Carlo”. To enhance performance we have used a variation of the Metropolis-Hastings prescription known as “Hybrid Monte Carlo”, that first appeared in the physics literature in 1987 [14]. A brief outline of the algorithm is sketched in Section 3 and we refer the reader to the appendix for a more extensive discussion.

For testing the performance of our proposed method, we use artificial data derived from known multivariate Gaussian distributions, calibrated using data from eight different asset classes for the last 14 years. This allows to simulate the finite-sample behaviour of our “best portfolio” estimator (PU from now on), and compare it to the standard MEU prescription. Since the real optimal portfolio - with respect to the chosen utility function - is known, the speed of convergence can be measured empirically. As will be shown in Section 4, our method clearly outperforms the simplistic optimisation. Interestingly one observes that up to a threshold of about 350 monthly observations (corresponding to almost 30 years of data) the knowledge gained from data is actually insufficient for selecting any but the uniformly distributed portfolio. We are also able to confirm a significant improvement with regard to the instability of the algorithm induced by the risk-aversion parameter. In the end some backtesting is performed: when looking at “what if” investment scenarios, our method again shows superior performance for at least one typical investment profile.

The paper is organised as follows: in Section 1 we propose our method, in which MEU optimisation is replaced by a double expectation with respect to (a transformation of) the utility function and the conditional posterior distribution. Section 2 is devoted to the analysis of the posterior distribution. In Section 3 we deal with numerical integration, and report empirical results in Section 4. Conclusions and final remarks are presented in Section 5.
1. The “Recommended” Portfolio Approach

After illustrating some typical features of the problem under examination by means of a simplified example, we will formally introduce our probabilistic interpretation of the utility function, followed then by the Bayesian analysis of the problem.

A simplified example

To illustrate what we said in the introduction, let us consider the following function:

\[ u(\alpha, M, \delta r) = (1 + \alpha \delta r) \left[ 1 + \exp \left( -\frac{\alpha}{M} \right) \right]^{-1} - \alpha^2 \left[ 1 + \exp \left( -\frac{\alpha - 1}{M} \right) \right]^{-1} \tag{1} \]

which is plotted twice in its dependence on \( \alpha \) in Fig.1. Both graphs correspond to a value of \( M = .01 \); they only differ in the choice of \( \delta r \) that is set to .01 and to -.01.

![Figure 1. Plot of the function \( u(\alpha) \) from Eq.1.](image)

With these minimal parameter modifications, the qualitative features of the two curves are virtually unchanged, but the maximum within the interval \( \alpha \in [0, 1] \) moves from \( \alpha_{\text{max}} = 0.91 \) to \( \alpha_{\text{max}} = 0.09 \). Clearly this represents a serious problem whenever the determination of parameters would be based on a fit to observed data which necessarily will be plagued by errors. The value for \( \delta r \) could, for example, be the expected excess return of our portfolio with respect to some low risk asset: a small error in \( \delta r \) would have a severe effect on the location of the maximum and, consequently, on the selected value for \( \alpha_{\text{max}} \). If the function \( u(\alpha) \) were to represent some sort of expected utility, it should be pointed out that just choosing a portfolio (mimicked here by the choice of a value for \( \alpha \)) by blind faith according to the maximising principle, would, for both values of \( \delta r \), lead to an unnecessary high amount of risk, since a minimal error in \( \delta r \)
could cause a deep fall respectively to the left or right of the two maxima. For both functions a more conservative settlement somewhere in the middle would not induce nearly as much risk and still achieve results not too different from those guaranteed by the apparent “optimal” choice.

In order to overcome this problem we suggest to interpret the expected utility as proportional to the logarithm of a probability in the space of portfolios, and replace the prescription of selecting the “optimum” portfolio as the maximum of the expected utility by the rule that the recommended portfolio is the expectation value of the portfolio weighted by its probability:

\[ E(\alpha|M, \delta r, \nu) = Z(M, \delta r, \nu) \int_0^1 d\alpha \alpha \exp(\nu u(\alpha, M, \delta r)), \]

(2)

where

\[ Z^{-1}(M, \delta r, \nu) \equiv \int_0^1 d\alpha \exp(\nu u(\alpha, M, \delta r)), \]

and \( \nu \) is a constant. This definition of \( E(\alpha|M, \delta r, \nu) \) is a continuous and differentiable function in all of its parameters as opposed to the discontinuity of the maximum prescription, and if we define

\[ \alpha^* = \lim_{\nu \to \infty} E(\alpha|M, \delta r, \nu), \]

it is easy to show that \( \alpha^* \) is the solution of

\[ \text{MAX}_\alpha u(\alpha, M, \delta r), \]

and we hence correctly recover the singular behaviour of maximisation as the limit value for perfectly analytic functions. The reader reluctant of replacing the maximisation principle can always interpret our prescription as a smooth interpolation procedure, much in the same spirit as the simulated annealing minimisation procedure introduced by Kirkpatrick et al [20]. In the remaining parts of this paper we will sometimes abuse the term “optimum” referring with it also to the portfolios produced by our approach. From the context it will always be clear what kind of optimum, MEU (maximising expected utility) or PU (probabilistic utility) we mean.

1.1. Probabilistic Interpretation of the Utility Function

Let us denote with \( \alpha \) the set of parameters that identify a portfolio, and with \( U \) the set of parameters that characterise our utility function model, like for instance risk aversion, investment horizon etc. Let us assume furthermore that the expected utility is computed with respect to a distribution characterised by parameters, like expected excess returns for instance, that we will denote collectively with \( \Phi \). The expected utility can then be written as a function

\[ u = u(\alpha, U, \Phi). \]

In classical asset allocation theory the prescription would be to select portfolios that maximize the expected utility; in our framework we decided to consider the expected
utility as proportional to the logarithm of a probability measure in portfolio space, fully conditional on $U$ and $\Phi$.

$$\alpha \sim P(\alpha|U, \Phi) = Z(\nu, U, \Phi) \exp \left( \nu u(\alpha, U, \Phi) \right).$$

(3)

The symbol $\sim$ has to be interpreted as: $\alpha$ is distribute according to, and $Z(\nu, U, \Phi)$ is a normalization constant defined by

$$Z^{-1}(\nu, U, \Phi) = \int_{D(\alpha)} [d\alpha] \exp \left( \nu u(\alpha, U, \Phi) \right),$$

where $D(\alpha)$ stands for the integration domain of $\alpha$.

The recommended portfolio $\overline{\alpha}$, given $U$ and $\Phi$, is defined as the expectation value of $\alpha$:

$$\overline{\alpha}(U, \Phi) = Z(\nu, U, \Phi) \int_{D(\alpha)} [d\alpha] \alpha \exp \left( \nu u(\alpha, U, \Phi) \right).$$

(4)

Since we choose to insist on a distribution to describe the portfolio, it is natural to identify the error associated with the estimate of $\overline{\alpha}$ with the standard deviation of the distribution itself. An unbiased estimate of the standard deviation will be computed, at no extra cost, while computing the integral in (3).

The parameter $\nu$ is a constant that the theory is unable to set. Its meaning though is quite direct. If we send $\nu \to 0$ we see that the density distribution for $\alpha$ becomes the uniform one, all portfolios are just as likely and the ideal one, according to the previous prescription, would just be evenly spread over all of the assets available. On the other hand, if we send $\nu \to \infty$, the ideal portfolio would just coincide with the one obtained by the standard MEU procedure. For an infinite $\nu$ all of the measure is just concentrated about the maximum of the expected utility. In short, $\nu$ measures the weight that expected utility should have as opposed to the total noise generated by the flat measure $[d\alpha]$. If we have a set of stationary historical data we can bootstrap from the data, build scenarios and compute unbiased estimates of the expected utility; or, if we have a data model and we believe that historical data are drawn from some distribution, we can use the time series to estimate the distribution parameters. In both situations our confidence on the value of the expected utility will be in some way linked positively to the length of the available time series data set. It seems a reasonable assumption for $\nu$ to exhibit the following asymptotic behaviour:

$$\lim_{N \to 0} \nu(N) = 0,$$

(5)

$$\lim_{N \to \infty} \nu(N) = \infty,$$

(6)

where $N$ is the size of the data set. The simplest such form is

$$\nu = \rho N^\gamma,$$

(7)

with $\rho$ and $\gamma$ constants strictly greater than 0. All of the simulations carried out in this paper will have $\rho = 1$ and $\gamma = 1$. The limit $\rho \to \infty$ will recover the standard maximisation approach. It is obviously interesting to ask whether a more sophisticated relation between $\nu$ and $N$ could lead to a better algorithm, in particular to one that makes a more effective use of the available information. However, since already the
simple link $\nu = N$ leads to great improvements, these questions will be addressed in future research.

1.2. Bayesian Analysis and Parameter Determination

Parameters that characterise the distribution of returns are determined with some degree of uncertainty that must be taken into account. A consistent framework to do so is to accept the Bayesian point of view that it is not possible to infer the values of model parameters from experimental data with certainty, and to think of parameters as random variables themselves, described by a distribution. Based on the observations, we modify our view in a consistent manner with the observed data. The result of this process will be a posterior distribution $P(\Phi|\{R\})$, i.e. a distribution fully conditional on the historical data $\{R\}$.

The uncertainty on the average returns must therefore play a role in the calculation of the optimised portfolio. The Bayesian prescription to do so is to replace Eq.4 with the following:

$$\alpha_{PU} = \int_{D(\Phi)} d\Phi \Pi(U, \Phi) P(\Phi|\{R\}).$$

To proceed with the computation of the integral on the r.h.s of Eq.8, we need to know the posterior distribution for $\Phi$ that from Bayes’ theorem turns out to be

$$P(\Phi|\{R\}) = \frac{P(\{R\}|\Phi) P_0(\Phi)}{P(\{R\})}.$$

The denominator $P(\{R\})$ is the unconditional distribution of the observed data $\{R\}$ and for our purposes but a normalisation constant, while the two terms in the numerator represent the more interesting ones. The quantity $P(\{R\}|\Phi)$ is the likelihood or probability density of the observed data subject to the fact that the parameters are exactly $\Phi$. The second term in the numerator of Bayes’ theorem, $P_0(\Phi)$, constitutes the a priori distribution for the parameters, embodying thereby personal views on the expected behaviour of the distribution of $\Phi$. A Bayesian approach requires you to state explicitly what theory underlies your assumption, and the place to do so is precisely in the choice of the prior $P_0(\Phi)$. A prior should be chosen in accordance to our knowledge and prejudices. If we have no reason to believe anything at all, the prior will reflect this by assigning equal probability to any possible configuration. It will become more and more decisive the stronger our convictions are rooted in background knowledge we have about the problem.

2. An Explicit Posterior Distribution

To proceed further we need to choose some particular data model. For the time being, and given the aims of this paper, we resort to a classical Gaussian framework. However, it is worth noting that the selection of the data model could be itself a subject of Bayesian inference: we defer this extension to future research. The posterior distribution of (8) can now be written out by data inspection. Denoting with $m$ the average returns
and with $\Omega$ the covariance matrix, the set $\{m, \Omega\}$ makes explicit what was previously referred to as $\Phi$.

The likelihood term of (9) can be written:

$$P(\{R\}|m, \Omega) = \prod_{n=1}^{N} \frac{\exp \left( -\frac{(r_n-m)^T \Omega^{-1} (r_n-m)}{2} \right)}{\sqrt{(2\pi)^J |\Omega|}}$$

$$= \exp \left( -\frac{N}{2} m^T \Omega^{-1} m + Nm^T \Omega^{-1} \bar{r} - \frac{1}{2} \sum_{n=1}^{N} r_n^T \Omega^{-1} r_n \right) \left[ \sqrt{(2\pi)^J |\Omega|} \right]^{-\frac{N}{2}},$$

(10)

where $J$ is the number of assets, $r_n$ the n-th observations vector, and:

$$\bar{r} = \frac{1}{N} \sum_{n=1}^{N} r_n.$$  

We will choose a prior for $m, \Omega$ of the form:

$$P_0(m, \Omega) = P_0(m|\Omega)P_0(\Omega),$$

(11)

where the average conditional distribution is chosen as a normal,

$$P_0(m|\Omega) \simeq \exp \left( -\frac{\beta}{2} (m - \chi)^T \Omega^{-1} (m - \chi) \right).$$

(12)

The vector $\chi$ is the view we hold, consistent with our background knowledge, of the central point of the distribution of average returns, while $\beta$ is a hyper-parameter that the theory cannot fix; it controls the width of the distribution and we will soon see a possible interpretation.

The prior for the covariance matrix is the inverse Wishart:

$$P_0(\Omega) \simeq |\Omega|^{-\frac{h+J+1}{2}} \exp \left( -\frac{1}{2} Tr \left[ \Omega^{-1} \Sigma_{\Omega} \right] \right),$$

(13)

where $h$ is once again another hyper-parameter and $\Sigma_{\Omega}$ is our view.

Putting all together, we have:

$$P_0(m, \Omega|\{R\}) = P_0(m|\Omega, \{R\})P_0(\Omega|\{R\}),$$

(14)

where:

$$P_0(m|\Omega, \{R\}) \simeq \exp \left( -\frac{N(1+\kappa)}{2} (m - \mathbf{M})^T \Omega^{-1} (m - \mathbf{M}) \right)$$

(15)

$$P_0(\Omega|\{R\}) \simeq |\Omega|^{-\frac{h+N+J+1}{2}} \exp \left( -\frac{N+h}{2} Tr \left[ \Omega^{-1} \mathbf{A} \right] \right)$$

(16)

$$\mathbf{M} = \frac{\bar{r} + \kappa \chi}{1 + \kappa},$$

(17)

$$\kappa = \frac{\beta}{N},$$

(18)

$$\mathbf{A} = \frac{h \Sigma_{\Omega}}{N+h} + \frac{N \Sigma}{N+h} + \frac{\kappa (\bar{r} - \chi) (\bar{r} - \chi)^T}{(1+\kappa)(N+h)}$$

(19)

$$\Sigma = \frac{1}{N} \sum_{n=1}^{N} (r_n - \bar{r}) (r_n - \bar{r})^T.$$  

(20)
All the details of the computation can be found in Appendix D. It seems natural to view $\kappa$ and $h$ as a simple way to measure the degree of confidence we have in our views as opposed to the indications stemming from historical data. If we hold a view but we think that observed data should weigh more in our decision process, then we would choose small values for $\kappa$ and $h$. Note that in the limit $\kappa \to 0$, we would recover, for the average return, a totally non-informative prior that assigns equal probability to any possible value of $m$. A strong view is represented by a large $\kappa$ and large $h$. In the limit $\kappa \to \infty$ and $h \to \infty$, the posterior distribution would be centred about our views regardless of the historical data, and the width of the distribution would tend to 0.

3. Numerical Integration

3.1. Markov Chain Monte Carlo Integration

The integral in Eq. (8) is easily carried out by Markov Chain Monte Carlo (MCMC) integration [16, 23]. Since the probability distribution for $(m, \Omega)$ is independent from the distribution of $\alpha$, an algorithm that generates the Markov Chain capable of yielding the correct distribution is as follows:

Step 1 Sample $\Omega$ from the inverse Wishart probability density function (p.d.f.):

$$|\Omega|^{\frac{k+N+J-1}{2}} \exp \left[ (N + h) \text{Tr}[\Omega^{-1} A] \right].$$

This can be achieved by generating $N + h$ $J$-dimensional arrays $x_i, i = 1, \ldots, N + h$ distributed according to

$$x \sim N(0, \Omega^{-1}),$$

and setting

$$A = \frac{1}{N + h} \sum_{i=1}^{N+h} x_i x_i^T.$$  

Step 2 Holding fixed the sampled $\Omega$, sample $m$ from the p.d.f.:

$$\exp \left( -\frac{N(1+\kappa)}{2} (m - M)^T \Omega^{-1} (m - M) \right).$$

Step 3 Holding fixed the values for $(m, \Omega)$, sample $\alpha$ from the p.d.f.

$$\exp \left( NE[u(\alpha, U, m, \Omega)] \right).$$

Details of the algorithm and the proof that produces an unbiased estimate of the integral in the r.h.s of Eq. (8) can be found in Appendix C.

Steps 1 and 2 do not present any problem since we know how to sample from those p.d.f.s. Step 3 is somewhat more complex. We do not know how to sample directly from that p.d.f., and we are forced to devise a Markov chain that relaxes to the desired distribution. After several experiments with variations of the Metropolis-Hastings, we resorted to an implementation of the "Hybrid Monte Carlo" method. Once
relaxation has been achieved we can run the Markov chain for few more steps in order to perform measurements. Relaxation or thermalisation is not a trivial issue but a thorough discussion of the problems involved would bring us too far from the subject of this paper. We choose to defer this discussion to a forthcoming paper focussing on the implementation of the numerical integration scheme.

3.2. The Utility Function

The selection of a good utility function is not the subject of this paper, nor is it particularly relevant for our results. Whenever the return distribution is assumed to be normal, as in our framework, the explicit solutions of all the utility functions are but a combination of first and second distribution moments. Still, a non-trivial difference arises when standard deviation terms are included, since they are able to generate a time horizon effect, i.e. an effect that favours less risky assets on the short range and turns on risky ones, with higher returns, on the long range. We are aware of the academic debate on this topic, testified by a considerable amount of related literature [10, 11, 21, 28, 29], and we believe this to be a desirable feature for a utility function. The probability for the riskier assets to outperform the less risky ones, in fact, approaches one asymptotically with time, being it the error function of the ratio between mean and standard deviation, which grows with the square root of time.

However, utility functions of standard use in financial economics (such as those who exhibit Constant Relative Risk Aversion) do not fall in this category. The standard deviation terms are directly related to non-regular utility functions that measure risk with the concepts of, say, Value at Risk, Loss Probability etc., i.e. with the so-called downside risk measures [8]. In this way risk is measured by the expected amount by which a specified target is not met: this might better describe how the investor perceives risk, as documented by results from behavioural finance [17], and is more in line with some recent ALM practice.

For these reasons we employed the following expected utility function, drawn from the article of Consiglio et al [13]:

\[
E \left[ u(\alpha, L, \lambda, T) \right] = \sum_{n=1}^{N_T} \Delta t [E(U(n\Delta t)) - \lambda E(D(n\Delta t))],
\]

(26)

where \( U(n\Delta t) \) and \( D(n\Delta t) \) are the upside and downside, respectively, of the portfolio return at time \( n\Delta t \) against a fixed target return \( L \), and \( \lambda \) is a weight indicating the investor risk aversion. The time horizon \( T \) is built out of \( N_T \) intermediate time intervals \( \Delta t \) such that \( T = N_T \Delta t \), is a sequence of \( N_T \) values for \( \omega(n\Delta t) \), \( n = 1, \ldots, N_T \). The model takes a "target-all time" view, and the allocation is such that staying as close to the target return trajectory at all times is the primary concern. A risk averse investor will want to keep as far as possible from target return under-performance situations, and will favour paths close to the target line.

Modelling the distributions for the single period log-return \( \omega \) with the normal
we obtain by straightforward (tedious) Gaussian integration an explicit expression for the utility function:

$$E\left[u(\alpha, L, \lambda, T)\right] = \sum_{n=1}^{N_T} \Delta t \left[ n\Delta t M f_2(n\Delta t) - \sqrt{n\Delta t} S f_1(n\Delta t) \right],$$

(28)

where:

$$f_1(t) = (\lambda - 1) \frac{e^{-t\eta^2}}{\sqrt{2\pi}}$$

(29)

$$f_2(t) = \frac{1 + \lambda}{2} - \frac{\lambda - 1}{2} \text{erfc}(\sqrt{t}\eta)$$

(30)

$$\eta = \frac{M}{2S}$$

(31)

$$M = \alpha^T \cdot m - L$$

(32)

$$S^2 = \alpha^T \cdot \Omega \cdot \alpha.$$  

(33)

As expected, the explicit solution for this specific form of utility is a function of the portfolio mean, variance and -most importantly- standard deviation. It incorporates a competing effect between average return and standard deviation with different time scaling properties: the standard deviation’s contribution is proportional to $\lambda - 1$, and can be traced back to the imperfect cancellation between positive and negative deviations from ideal line. We thus obtain the desired dependency of the optimal portfolio on the chosen time horizon: the longer the horizon (ceteris paribus), the more aggressive the optimal allocation.

4. Empirical Results

In this section the performance of our proposed PU method will be analysed in various contexts: first its consistency and speed of convergence will be tested and compared to the MEU optimisation with the help of artificial data generated by a known multivariate normal distribution. Afterwards the performance of both prescriptions will be reviewed by means of historical time series data. As mentioned before, the sensitiveness towards the risk aversion parameter $\lambda$ of both selection procedures will also be evaluated. Finally, the effect of incorporated personal views is illustrated, and the degree of confidence associated with an “optimised” portfolio is discussed.

Historical data used to infer distribution parameters consists of 8 monthly indexes covering the period from January 1988 to January 2002. In Tabl.1 we show the list of titles employed; this set of data will be referred to as full sample in the following.
Table 1. List of assets employed. The full set of the data goes from Jan 1988, to Jan 2002. The used acronyms have the following meaning: MSCI = Morgan Stanley Capital Index, JPM = JPMorgan Index, ML = Merrill Lynch Index. Data source: Datastream. Data types: Price Index for equities, Total Return Index for bonds. All the samples are in local currency, unadjusted for inflation. The index titles refer to the Datastream mnemonics.

| Assets     | Description                      |
|------------|----------------------------------|
| MSNAMR     | MSCI North America Equity        |
| MSPACF     | MSCI Pacific Equity              |
| MSEROP     | MSCI Europe Equity               |
| JPMUSU     | JPM US Government Bond           |
| JPMJPU     | JPM Japan Government Bond        |
| JPEMEL     | JPM Europe Government Bond       |
| MLHMAU     | ML US Corporate High Yield       |
| JPEC3M     | JPM Euro Cash                    |

4.1. Simulation with Artificial Data

We first investigate the performance of our proposed PU method by using artificial data to simulate finite sample behaviour. For the testing we assume the true return distribution to be a multivariate Gaussian, characterised by the parameters estimated from the full historical sample. From this distribution we generate 1000 independent samples of various fixed lengths. For any given sample length and a fixed parameter set (L=5% per yr, T=1 yr, and λ = 3), we then calculate the average Euclidean distance of both the MEU and PU 1000 optimal portfolios from the “truly” optimal allocation, that we can determine exactly from the parameters of the assumed “true” distribution. In Fig.2 we have plotted the results of this exercise, together with a straight line showing the average distance of a randomly chosen portfolio from the “true” one. Our PU method clearly outperforms the MEU procedure, for it is always closer to the true allocation and below the random-choice threshold. The picture well illustrates the extreme sensitiveness of the MEU procedure to the input data; for a great distance from a benchmark portfolio when averaging over 1000 samples can only be explained by a great variability in the portfolio composition over the different samples.

Asymptotically, for $N \to \infty$, the return distribution parameters are determined with quasi-certainty, and we consequently recover the “true” optimal portfolio, thereby verifying the consistency of both approaches. In Table 2 we present evidence for this, reporting the allocations for $N = 32000$ observations.

However, for the classical MEU optimisation the speed of convergence looks worryingly slow when considering typical lengths of time series data used in asset allocations by practitioners. Indeed, for the chosen set of parameters one observes that up to a threshold of more than 350 monthly observations, corresponding to a data sample of almost 30 years, the knowledge gained from data is actually insufficient for selecting

† for a derivation of this value refer to Appendix A
any but the equally-weighted portfolio! This nicely illustrates the real risk of estimation errors completely overwhelming the value of information contained in the data. A restriction to very long data sets could seem a solution (provided data is available), but then one could object again by referring to the well known non-stationarity exhibited by financial time series. On the other hand, our PU prescription manages to stay always below the random portfolio threshold line, although coming very close to it when observations are scarce, thereby justly reflecting a situation in which data is not sufficient to justify very “particular” portfolios.

**Figure 2.** Distance from TRUE Portfolio w.r.t. sample size: for each sample length, average euclidean distance of the MEU and PU portfolios (resulting from 1000 independent samples) from the known true optimal allocation. The benchmark value for a randomly chosen portfolio is represented by the straight line.

| ASSET       | PU   | MEU |
|-------------|------|-----|
| MSNAMR      | 0.04 ± 0.01 | 0.04 |
| MSPACF      | 0.00 ± 0.00 | 0.00 |
| MSEROP      | 0.00 ± 0.00 | 0.00 |
| JPMUSU      | 0.02 ± 0.02 | 0.02 |
| JPMJPJ      | 0.00 ± 0.00 | 0.00 |
| JPMEIL      | 0.28 ± 0.08 | 0.27 |
| MLHMAU      | 0.04 ± 0.02 | 0.04 |
| JPEC3M      | 0.62 ± 0.10 | 0.63 |

**Table 2.** Optimal portfolio for N=32000. The investment parameters are constant and set to $\lambda = 3$, $T=1$ yr, $L=5\%$. 
4.2. Backtesting on Historical Data

Based on the full sample historical data and for some chosen set of parameters, we have performed some backtesting, in the form of “what if” investment scenarios. To this end, we used 5 years rolling windows for estimation and 3 years rolling windows for out-of-sample testing, together with two larger samples. We measure each portfolios’ hypothetically achieved performance. In Table 3 we present back testing results for different samples; for each sample, we have selected an “average” risk attitude, and computed a unique performance indicator (Sharpe Ratio), neglecting the behaviour at intermediate intervals. Results vary over the examined samples: until the first half of the ’90s, when financial series exhibited more stable patterns, the MEU procedure achieves better performances, while for more recent samples it is the PU that outperforms the MEU optimisation. Using longer samples advantages the PU procedure.

| Estimation Sample | Out-of-Sample | Sharpe Ratio |
|-------------------|---------------|--------------|
| 1988 – 1992       | 1993 – 1996   | 3.17         |
| 1989 – 1993       | 1994 – 1997   | 2.64         |
| 1990 – 1994       | 1995 – 1998   | 2.63         |
| 1991 – 1995       | 1996 – 1999   | 2.10         |
| 1992 – 1996       | 1997 – 2000   | 1.61         |
| 1993 – 1997       | 1998 – 2001   | 0.84         |
| 1994 – 1998       | 1999 – 2002   | –0.23        |
| 1988 – 1995       | 1996 – 2002   | 0.71         |
| 1988 – 1997       | 1998 – 2002   | 0.20         |

Table 3. Back-testing.

4.3. Sensitiveness Towards Risk Aversion Parameter \( \lambda \)

As a second empirical investigation, we examine the algorithms’ stability for a given portfolio profile (\( L=5\% \) per yr, \( T=1 \) yr). As previously stressed, all expected-utility based procedures suffer from the presence of a risk aversion parameter, dimensionless and un-settable from theory. As a measure of instability, it seems then natural to compare the sensitivity to \( \lambda \) for both the MEU optimisation prescription and the PU method we propose in this paper. Specifically, we examine the behaviour of a diversification indicator, i.e. an indicator that measures the degree of concentration within a portfolio, and consequently allows to identify the range of the parameter that mostly affects the portfolio composition.

The simplest of such a - as Bouchaud et al. put it - entropy-like measure is the

\( \dagger \) Of course, such an ex post performance verification for some (by us) chosen set of time series and investor profile does not allow to draw definite conclusions; it is merely meant to support and illustrate the more important results from the above section.
quantity:
\[ Y = \sum_{j=1}^{J} \alpha_j^2. \]  

which ranges from \( \frac{1}{J} \) (\( J \)=number of assets= 8 here), when the portfolio is totally diversified (evenly spread), to 1 in case of complete concentration on one asset.

In Fig.3 and Fig.4 we report the behaviour of \( Y \) with respect to \( \lambda \) for two different data samples, the full sample and a slightly restricted 1988-2000 one. Looking at the MEU graph in Fig.3, the behaviour appears very erratic and the significant range of \( \lambda \) restricted to a relatively small interval, meaning that small changes in \( \lambda \) can produce large modifications in the portfolio composition. Indeed, if we look at Table 4, we can observe how the portfolio composition changes as \( \lambda \) moves from 2.6 to 3.0, to the point that the portfolios are totally twisted around. This is certainly not reassuring, given that \( \lambda \) is only loosely tied to investor’s risk aversion, and its setting is not without uncertainties. Back to Fig.3, what strikes even more is what happens when we look at the results for a different data sample, in this case shortened by the last two years of observations: the curve decidedly shifts to the right, and consequently the relevant range of \( \lambda \) does the same, leading to dangerous risk profile mis-identifications, and forcing to re-calibrate (with all the associated uncertainties) the values of \( \lambda \) basically each time new historical observations are added to the sample.

Coming now to the PU model, for which results are shown in Fig.4, the diversification indicator displays a very different pattern: it indicates a more conservative overall behaviour, with values closer to the lower bound of \( \frac{1}{8} \). It never concentrates all the weights on a single asset, not even for the risk-neutral (\( \lambda = 1 \)) case. In Table 4 we can see data from our Bayesian PU approach: the variations in the portfolios induced by the different \( \lambda \)'s are now hardly noticeable. Most importantly, \( Y \) exhibits a smooth pattern. This reduces the danger of mis-settings of \( \lambda \) and its sensitiveness on the chosen sample; for different data sample, in fact, the curve shifts but remains rather similar, leaving unaffected the significant range of \( \lambda \).

| ASSET   | MEU | PU |
|---------|-----|----|
|         | \( \Lambda_1 \) | \( \Lambda_2 \) | \( \Lambda_3 \) | \( \Lambda_1 \) | \( \Lambda_2 \) | \( \Lambda_3 \) |
| MSNAMR  | 0.11 | 0.06 | 0.04 | 0.16 ± 0.25 | 0.14 ± 0.23 | 0.13 ± 0.22 |
| MSPACF  | 0.00 | 0.00 | 0.00 | 0.01 ± 0.01 | 0.01 ± 0.01 | 0.01 ± 0.01 |
| MSEROP  | 0.01 | 0.00 | 0.00 | 0.08 ± 0.15 | 0.07 ± 0.14 | 0.06 ± 0.12 |
| JPMUSU  | 0.00 | 0.02 | 0.00 | 0.15 ± 0.19 | 0.15 ± 0.18 | 0.14 ± 0.18 |
| JPMJPJ  | 0.00 | 0.02 | 0.00 | 0.03 ± 0.04 | 0.03 ± 0.04 | 0.03 ± 0.04 |
| JPMIEIL | 0.10 | 0.04 | 0.00 | 0.15 ± 0.20 | 0.14 ± 0.19 | 0.13 ± 0.18 |
| MLHMAU  | 0.00 | 0.43 | 0.28 | 0.20 ± 0.26 | 0.23 ± 0.23 | 0.23 ± 0.23 |
| JPEC3M  | 0.00 | 0.43 | 0.64 | 0.20 ± 0.26 | 0.23 ± 0.28 | 0.26 ± 0.30 |

Table 4. Optimal portfolio with respect to \( \lambda \). For all columns the time horizon \( T \) is one year and expected return 5% per year. The whole sample (1988-2002) is considered. The parameter \( \lambda \) instead is set to: \( \Lambda_1 : \lambda = 2.6 \), \( \Lambda_2 : \lambda = 2.8 \) and \( \Lambda_3 : \lambda = 3 \).
4.4. Confidence Associated with Identified Portfolio

While the MEU optimisation procedure places infinite faith on the distribution as determined through simplistic inspection of historical data, and makes no allowance for imperfect knowledge, the Bayesian PU approach has this naturally built in. The
decidedly more conservative allocations that are manifested in our more balanced portfolios reflect the fact that we do not exactly know what the true distribution is, and thus we try to protect ourselves against situations in which actual distributions are rather different from the ones we are invited to deduce from historical data. Another hint for the relative smallness of the full sample comes from the observation of the portfolio’s standard errors quoted in Tabl.4. They are of the same magnitude of the average value (=weight) of the asset, indicating that we should not take too seriously a prediction of 14% as opposed to a 20%. From this table we can safely conclude which assets should not be in our portfolio, while when it comes to the single best way to distribute the others we can, at best, be only suggestive. The confidence intervals might appear too large, but they are just another confirmation ex post of the inner consistency of our formalism: from Fig.2 we know that the square distance of our estimated portfolio from the ”true” portfolio should be of the order of \( d^2 = 0.57^2 = 0.32 \) in this specific case (full sample, 180 observations). As a rough test, we might double check this number by summing the squares of the PU calculated standard errors as displayed in Tabl.4, where we find 0.27, which results very well compatible within this rough consistency check.

Since MEU optimisation completely trusts its return distribution parameters as estimated by available data, it naturally misses a means to characterise the degree of confidence to be attached to its ”optimal” portfolio. However, from our finite sample tests as shown in Fig.2 we are able to give a rough estimate of the mean standard error of every asset’s weight: 0.77 is the average distance from the true portfolio in the case of \( N = 180 \) (our full sample), and therefore \( 0.77/\sqrt{8} = 0.27 \) should be a reasonable estimate of the error, ignored by the MEU formalism, but definitely a reality which should not be denied.

4.5. Effects of Incorporated Personal Views

In Tabl.5 it is illustrated what happens when we express personal views for the distribution moments. In the first column we report the portfolio allocation for neutral views. In the second one, we have incorporated views only on the mean values of the equity indexes: we postulate a very optimistic scenario, with an annual average return of 11% for MSNAMR and MSEROP, and of 15% for MSPACF. We attach a rather strong degree of confidence to this personal view, setting \( k = 10 \) (i.e. \( \beta = 10N \)). The results are in line with the previously expressed views: MSPACF, that was not selected in the neutral-views scenario because of the poor performance over our historical data sample, is now the most over-weighted asset class, since it was modified by our strong expectation. The portfolio errors drop consequently, because of the confidence degree attached to the views. In column three we repeat the exercise with views on variances. We express views, again only on equities, based on the implied volatilities inferred from proxy indexes options with two years expiration. The values are quite large if compared to the historical ones (annualised implied volatilities: MSNAMR 25% MSPACF 26%, and MSEROP 29%), reflecting the market sentiment for the near future. We set \( h = N \), so that the resulting variances are the average between the historical and the implied
ones. As expected, the resulting asset allocation is more conservative, and, given the errors size, almost all asset classes are included in the portfolio.

| ASSET     | PU                      |
|-----------|-------------------------|
|           | $k = 0, h = 0$ | $k = 10, h = 0$ | $k = 0, h = N$ |
| MSNAMR    | 0.39 ± 0.41           | 0.23 ± 0.24           | 0.20 ± 0.31           |
| MSPACF    | 0.00 ± 0.03           | 0.58 ± 0.30           | 0.06 ± 0.16           |
| MSEROP    | 0.19 ± 0.34           | 0.07 ± 0.13           | 0.15 ± 0.26           |
| JPMUSU    | 0.08 ± 0.22           | 0.01 ± 0.05           | 0.12 ± 0.24           |
| JPMJPJ    | 0.00 ± 0.01           | 0.00 ± 0.00           | 0.07 ± 0.18           |
| JPMEIL    | 0.18 ± 0.32           | 0.08 ± 0.17           | 0.15 ± 0.26           |
| MLHMAU    | 0.15 ± 0.31           | 0.03 ± 0.10           | 0.16 ± 0.29           |
| JPEC3M    | 0.01 ± 0.06           | 0.00 ± 0.00           | 0.10 ± 0.25           |

Table 5. Incorporating views: portfolio selection for neutral views (column 1), views on means (column 2) and views on variances (columns 3). The parameters set are constant and equal to $\lambda = 3, T = 5\text{years}, L = 11\%$.

5. Conclusion and final remarks

The purpose of this work was to address and improve some of the well known weaknesses of portfolio selection by maximising expected utility. We have pointed out that seeking and settling on an extremum of a utility function is equivalent to claiming absolute knowledge of the parameters governing the distribution of average returns. While theoretically this is never the case, historical data might at best offer partial support to our selection process, for which we attempted to provide a unified framework. The approach presented here takes into account parameter uncertainty and greatly reduces the instability of results common in standard optimisation procedures. We achieved this by employing a different interpretation of the utility function, and by endorsing a Bayesian framework approach. In doing so one benefits from several advantages: the framework provides a consistent way to account for uncertainty and, whenever we hold views, we can readily introduce them. Moreover, the standard error calculated easily for the recommended portfolio gives a good idea about the degree of certainty offered by the available historical data. We have tested the proposed method against traditional expected utility maximisation, using artificial data to simulate finite-sample behaviour, and have shown superior performance of our method as compared to the simplistic optimisation. This picture was reinforced when backtesting with historical data. We also managed to significantly improve the intrinsic instability with respect to the risk-aversion parameter (lack of continuity) that plagues all maximisation approaches.

As for future lines of research, we might be interested in relaxing the normality assumption, for instance by modelling the data with a mixture of Gaussian distributions: in Section 2 we hinted that the selection of the data model could itself be a subject of Bayesian inference. Additionally, there were some occasions in which our theory led
to parameters or hyper-parameters that could easily be determined in their asymptotic
behaviour, but whose value on the intermediate range was not clear (ν in Eq.4, h
in Eq.13, β in Eq.12). Especially ν, the smoothing parameter within our probabilistic
utility certainly has an important influence on the overall performance of our method; it
would therefore be interesting to ask whether a more sophisticated prescription than the
by us employed ν = N could lead to an enhanced overall performance, i.e., in particular,
to a faster convergence towards any “true” optimal portfolio.

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Appendix A. Random Portfolios

Let us denote with \( \mathbf{p} \) a particular portfolio. It is instructive to ask what would be the average square distance from it if we were to draw random portfolios. In this context 'random' means portfolios drawn uniformly from the hyperplane characterised by the equality constraint:

\[
\sum_{j=1}^{J} p_j = 1
\]

and the \( J \) inequality constraints

\[
p_i \geq 0,
\]

\[
\ldots,
\]

\[
p_j \geq 0.
\]

The expected value of a function with respect to this measure is defined as:

\[
E[O] \equiv Z \int_0^1 dp_1 \int_0^{1-p_1} dp_2 \ldots \int_0^{1-\sum_{j=1}^{J-2} p_j} dp_{J-1} O(p_1, \ldots, p_{J-1}, 1 - \sum_{j=1}^{J-1} p_j),
\]

where \( Z \) is a normalisation constant defined by

\[
1 = Z \int_0^1 dp_1 \int_0^{1-p_1} dp_2 \ldots \int_0^{1-\sum_{j=1}^{J-2} p_j} dp_{J-1}.
\]

Recalling the result

\[
\int_0^1 dp_1 \int_0^{1-p_1} dp_2 \ldots \int_0^{1-\sum_{j=1}^{J-2} p_j} dp_{J-1} p_1^{a_1} \ldots p_{J-1}^{a_{J-1}} = \frac{\Pi_{j=1}^J \Gamma(1 + a_j)}{\Gamma(\sum_{j=1}^J (1 + a_j))},
\]

we have:

\[
Z = \Gamma(J)
\]

\[
E[p_i] = \frac{1}{J}
\]

\[
E[p_i p_j] = \frac{1}{J(J + 1)} \quad \text{for} \quad i \neq j
\]

\[
E[p_i^2] = \frac{2}{J(J + 1)}
\]

and the average square distance is given by:

\[
E[\sum_{j=1}^{J} (p_j - \mathbf{p}_j)^2] = \sum_{j=1}^{J} \mathbf{p}_j^2 - \frac{2}{J(J + 1)}.
\]

Appendix B. Notation and detailed balance

In this appendix we introduce the basic concepts of Markov Chain Monte Carlo for the sole purpose of reviewing notation and fundamental results. The field is too vast to get into any depth within a few pages. The interested reader might refer to the literature on the subject, like for instance Gilks, Richardson and Spiegelhalter [16], and references cited therein.
Appendix B.1. Monte Carlo Integration

Let \( P(x) \) be a probability density for a random variable \( x \). If we draw samples \( \{x_i, i = 1, \ldots, n\} \) from \( P(x) \), we can evaluate the average \( E[g(x)] \) of an arbitrary function \( g(x) \)

\[
E[g(x)] \approx \frac{1}{n} \sum_{i=1}^{n} g(x_i). \tag{B.1}
\]

Appendix B.2. Markov Chains

Let \( T(x|y) \) a matrix describing the probability to get 'x' if we have 'y', then we can generate a Markov Chain (sequence of random variables) \( x_1, x_2, \ldots, x_t, \ldots \) such that the probability to get \( x_t \) is described by \( T(x_t|x_{t-1}) \).

If \( T(x|y) \) satisfies the equation:

\[
P(x) = \int dy T(x|y) P(y), \tag{B.2}
\]

the following theorem holds:

**Theorem 1:** Let \( \{x_0, x_1, \ldots, x_t, \ldots\} \) be the Markov Chain generated with transition probability \( T(x|y) \). If \( T(x|y) \) satisfies equation (B.2), with \( P(x) \) a given probability distribution, then uniform (unbiased) sampling from \( \{x_0, x_1, \ldots, x_t, \ldots\} \), will yield \( x_i \) with probability \( P(x_i) \).

Under these condition the probability \( P(x) \) is said to be the equilibrium distribution or the stationary point for \( T(x|y) \).

Appendix B.3. Markov Chain Monte Carlo Integration

From Theorem (1) and Eq. (B.1) it follows immediately that from any subsequence of a Markov Chain we can get an unbiased estimation of a function average, that is:

\[
\lim_{T \to \infty} \frac{1}{T} \sum_{t=t_s}^{t_s+T} g(x_t) = E[g(x)]. \tag{B.3}
\]

As it turns out, most of the time it is quite impossible to sample directly from a given distribution, but it is remarkably simple to create a Markov Chain that admits that same distribution as its stationary point.

Appendix B.4. Detailed Balance

The transition probability \( T(x|y) \) is a real probability in \( x \), that is

\[
\int dx T(x|y) = 1,
\]

and we can easily see that a sufficient condition for Eq. (B.2) to hold is to have:

\[
T(x|y)P(y) = T(y|x)P(x). \tag{B.4}
\]

This equation is called the detailed balance equation. From detailed balance, equation (B.2) follows directly after integrating in \( y \) both sides of Eq. (B.4).
The desired $T(y|x)$ can be built in virtue of the following:

**Theorem 2:** Let $\alpha(y|x)$ be any transition probability, then $P(y)$ will be a stationary distribution for $T(y|x)$ if

$$T(y|x) = \min\left(1, \frac{\alpha(x|y)P(y)}{\alpha(y|x)P(x)}\right)\alpha(y|x). \hspace{1cm} (B.5)$$

A particularly simple condition of application of this theorem is when $\alpha(x|y) = \alpha(y|x)$, in which case the prescription to build the transition matrix becomes:

- from a point $x_t$ propose a new point $y$ with probability $\alpha(y|x_t)$;
- if $P(y) > P(x_t)$ then set $x_{t+1} = y$, otherwise with probability $P(y)/P(x_t)$ set $x_{t+1} = x_t$.

### Appendix C. MCMC for portfolio optimisation

If we choose the transition probability:

$$\alpha(p_{t+1}, m_{t+1}|p_t, m_t) = P_p(p_{t+1}|m_{t+1})P_0(m_{t+1})$$

we have, according to Eq: $(B.3)$,

$$T(p_{t+1}, m_{t+1}|p_t, m_t) = \alpha(p_{t+1}, m_{t+1}|p_t, m_t).$$

Such a transition is readily obtained by sampling $m_{t+1}$ from the distribution $P_0(m)$, then, holding fixed $m_{t+1}$, sampling $p_{t+1}$ from the full conditional $P_p(p|m)$.

Sampling from $P_0(m)$ offers no challenge given that the random variable is normally distributed; the whole challenge is sampling $p$ from its fully conditional distribution.

This can be done by devising a suitable Markov chain with stationary distribution $P(p|m)$.

#### Appendix C.1. Metropolis MCMC

The first algorithm we present is a very simple implementation of the evergreen Metropolis algorithm.

From a location $p_t$, generate a random vector $v_t$ and consider the point

$$q = p_t + \epsilon v_t,$$  \hspace{1cm} (C.1)

where $\epsilon$ is a small number. Let

$$\Delta U \equiv E[u(q, L, \lambda, T)|m] - E[u(p, L, \lambda, T)|m],$$

and with probability

$$\pi = \min\left(1, \exp\left(\frac{\Delta U}{N}\right)\right)$$  \hspace{1cm} (C.2)

set $p_{t+1} = q$, and with probability $1 - \pi$, set $p_{t+1} = p_t$.

The step described in equation $(C.1)$ guarantees that the transition probability for the process $p_t \rightarrow p_{t+1}$ is the same as the transition probability for the inverse process $p_{t+1} \rightarrow p_t$. This suffices to prove that the Markov chain has the desired equilibrium
distribution. The only warning to be issued concerns the range of the variables \( p \). The domain \( D(p) \) is bounded therefore it will happen that step (C.4) will try to get on the outside. In this case care must be taken to bounce properly (a billiard ball rule will suffice) the trajectory in order to keep the point inside the domain.

**Appendix C.2. Hamiltonian MCMC**

The second algorithm we present is well known in the physics literature with the name of hybrid Monte-Carlo ([14]). In this appendix we limit ourselves to a short introduction.

Since we have to sample \( p \) keeping \( m \) fixed we are only interested in the functional form on \( p \) of the full conditional \( P_p(p|m) \):

\[
P_p(p|m) \sim \exp(U(p)).
\]

Expectations of functions of \( p \) will not be affected if we replace \( P_p(p|m) \) with the distribution

\[
G(p, \pi|m) \sim \exp\left(U(p) - \frac{\pi \cdot \pi}{2}\right), \tag{C.3}
\]

then starting from a pair \((p_n, \pi_n)\), the updating rule is defined as follows:

**Step 1** Sample \( \eta \) as a normal variable with mean zero and variance 1.

**Step 2** For a time interval \( T \), integrate Hamilton's equations

\[
\begin{align*}
\frac{d\pi_i}{dt} &= -\frac{\partial U}{\partial \pi_i} \tag{C.4} \\
\frac{dp_i}{dt} &= \pi_i, \tag{C.5}
\end{align*}
\]

together with the boundary conditions

\[
\begin{align*}
\pi(0) &= \eta, \tag{C.6} \\
p(0) &= p_n; \tag{C.7}
\end{align*}
\]

**Step 3** With probability

\[
\beta = \min\left(1, \exp(G(p(T), \pi(T)) - G(p_n, \eta))\right), \tag{C.8}
\]

set \( p_{n+1} = p(T) \), and with probability \( 1 - \beta \) set \( p_{n+1} = p_n \).

The clever idea behind this algorithm rests on the observation that, if step 2 is carried out exactly, Hamilton’s equation enforce \( G(p(T), \pi(T)) = G(p_n, \eta) \) and every proposed configuration is accepted. In general the acceptance rate will be controlled by the numerical accuracy of our integration scheme. A good scheme is the interleaved leap frog that, for finite integration step \( \Delta t \) and fixed trajectory length (that is, scaling the number of steps in the integration scheme with \( 1/\Delta t \)), is guaranteed to have errors \( O(\Delta t^2) \).
Appendix D. Likelihoods, Priors and Posteriors

Appendix D.1. Likelihoods of Data

The likelihood of observed data or the conditional density of data w.r.t a given model \( \{m, \Omega\} \) is given by:

\[
L(\{R\}|m, \Omega) = \prod_{n=1}^{N} \exp \left( -\frac{1}{2} (r_n - m)^T \Omega^{-1} (r_n - m) \right) \frac{d\nu(r_n)}{\sqrt{(2\pi)^J}|\Omega|} 
\]

\[
= \exp \left( -\frac{1}{2} \sum_{n=1}^{N} (r_n - m)^T \Omega^{-1} (r_n - m) \right) \prod_{n=1}^{N} \frac{d\nu(r_n)}{\sqrt{(2\pi)^J}|\Omega|} 
\]

\[
= \exp \left( -\frac{N}{2} m^T \Omega^{-1} m + Nm\Omega^{-1}\bar{r} - \frac{1}{2} \sum_{n=1}^{N} r_n^T \Omega^{-1} r_n \right) \prod_{n=1}^{N} \frac{d\nu(r_n)}{\sqrt{(2\pi)^J}|\Omega|} 
\]

where

\[
\bar{r} = (1/N) \sum_{n=1}^{N} r_n 
\]

The exponent can be written as:

\[
-\frac{N}{2} (m - \bar{r})^T \Omega^{-1} (m - \bar{r}) - \frac{1}{2} \sum_{n=1}^{N} r_n^T \Omega^{-1} r_n = 
\]

\[
-\frac{N}{2} (m - \bar{r})^T \Omega^{-1} (m - \bar{r}) - \frac{1}{2} \sum_{n=1}^{N} (r_n - \bar{r})^T \Omega^{-1} (r_n - \bar{r}) = 
\]

\[
-\frac{N}{2} (m - \bar{r})^T \Omega^{-1} (m - \bar{r}) - \frac{N}{2} Tr \left[ \Omega^{-1} \Sigma \right] 
\]

where \( \Sigma \) is the symmetric matrix whose element \( \{ij\} \) is:

\[
\Sigma_{ij} = \frac{1}{N} \sum_{n=1}^{N} (r_n - \bar{r})_i (r_n - \bar{r})_j 
\]

Appendix D.2. Priors

The prior for the model \( \{m, \Omega\} \) is given by:

\[
\Pi_0(m, \Omega|I) = \Pi_m(m|\Omega, I)\Pi_{\Omega}(\Omega|I) \quad (D.3) 
\]

where:

\[
\Pi_m(m|I) = \exp \left( -\frac{\beta}{2} (m - \chi)^T \Omega^{-1} (m - \chi) \right) \frac{d\nu(m)}{\sqrt{(2\pi)^J}|\Omega|} 
\]

and

\[
\Pi_{\Omega}(\Omega|I) = K|\Omega|^{-\frac{h+1}{2}} \exp \left( -\frac{h}{2} Tr \left[ \Omega^{-1} \Sigma \right] \right) d\mu(\Omega), 
\]

with \( K \) a constant of proportionality independent of \( \Omega \).
The measure $d\nu(x)$ is a measure in $\mathbb{R}^J$ while $d\mu(\Omega)$ is a measure in the space of symmetric positive definite matrices. Since any symmetric positive definite matrix $\Omega$ admits a unique decomposition

$$\Omega = O^T \Lambda O,$$

where $O$ is an orthogonal matrix in $J$ dimensions and

$$\Lambda_{ij} = \lambda_j \delta_{ij},$$

a diagonal positive definite matrix, the measure $d\mu(\Omega)$ decomposes in

$$d\mu(\Omega) = d\nu_+(\lambda) \left[ O^T dO \right],$$

where $d\nu_+(\lambda)$ is the flat measure in the semisphere $\mathbb{R}^J_+$ and $\left[ O^T dO \right]$ is the Haar measure on the orthogonal group in $J$ dimensions.

### Appendix D.3. Marginal Distributions

The marginal distribution for the observed data $M(\{R\}|I)$ is given by:

$$\int_{m,\Omega} L(\{R\}|m, \Omega, I) \Pi_0(m, \Omega|I) =$$

$$K \prod_{n=1}^N \frac{d\nu(r_n)}{\sqrt{(2\pi)^J}} \int_{m,\Omega} \frac{d\nu(m)}{\sqrt{(2\pi)^J}} d\mu(\Omega) \left| \Omega \right| \frac{N+1}{2} \exp \left( H(m, \Omega) \right) \quad (D.4)$$

where

$$H(m, \Omega) = -\frac{N}{2} (m - \bar{r})^T \Omega^{-1} (m - \bar{r}) - \frac{\beta}{2} (m - \chi)^T \Omega^{-1} (m - \chi)$$

$$- \frac{N}{2} Tr \left[ \Omega^{-1} \Sigma \right] - \frac{h}{2} Tr \left[ \Omega^{-1} \Sigma \Omega \right]$$

$$= - \frac{N + \beta}{2} m^T \Omega^{-1} m + N m^T \Omega^{-1} (\bar{r} + \kappa \chi)$$

$$- \frac{N}{2} \bar{r}^T \Omega^{-1} \bar{r} - \frac{\beta}{2} \chi^T \Omega^{-1} \chi$$

$$- \frac{N + h}{2} Tr \left[ \Omega^{-1} \left( \frac{h \Sigma \Omega}{N + h} + \frac{N \Sigma}{N + h} \right) \right]$$

$$= - \frac{N(1 + \kappa)}{2} (m - M)^T \Omega^{-1} (m - M)$$

$$- \frac{N}{2} Tr \left[ \Omega^{-1} \left( \bar{r} \bar{r}^T - (1 + \kappa) MM^T + \kappa \chi \chi^T \right) \right]$$

$$- \frac{N + h}{2} Tr \left[ \Omega^{-1} \left( \frac{h \Sigma \Omega}{N + h} + \frac{N \Sigma}{N + h} \right) \right]$$

where

$$H = \frac{\beta}{N}$$

$$M = \frac{\bar{r} + \kappa \chi}{1 + \kappa}. \quad (D.8)$$

$$\kappa = \frac{\beta}{N}. \quad (D.9)$$
If we define the matrix
\[ C = r r^T + \kappa \chi \chi^T - (1 + \kappa) M M^T \]
we get:
\[
C = \frac{1}{1 + \kappa} \left( (1 + \kappa) r r^T + \kappa (1 + \kappa) \chi \chi^T - (\bar{r} + \kappa \chi)(\bar{r} + \kappa \chi)^T \right) \quad (D.10)
\]
\[
= \frac{1}{1 + \kappa} \left( \kappa r r^T + \kappa \chi \chi^T - \kappa \bar{r} \chi^T - \kappa \chi \bar{r}^T \right) \quad (D.11)
\]
\[
= \frac{\kappa}{1 + \kappa} (\bar{r} - \chi)(\bar{r} - \chi)^T \quad (D.12)
\]
by which we get the final expression:
\[
H(m, \Omega) = -\frac{N(1 + \kappa)}{2} (m - M)^T \Omega^{-1} (m - M) - \frac{N + h}{2} T r \left[ \Omega^{-1} A \right], \quad (D.13)
\]
where the matrix \( A \) is given by:
\[
A = \frac{h \Sigma_{\Omega}}{N + h} + \frac{N \Sigma}{N + h} + \frac{\kappa (\bar{r} - \chi)(\bar{r} - \chi)^T}{(1 + \kappa)(N + h)}. \quad (D.14)
\]
The posterior is characterized by the following:
\[
m|\Omega \sim N(\bar{r} + \kappa \chi, \Omega) \quad (D.15)
\]
\[
\Omega \quad \sim W^{-1}(N + h, A). \quad (D.16)
\]

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