

# Risk and performance optimization for portfolios of bonds and stocks

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## Abstract

We explain how to optimize portfolios of bonds and stocks with respect to the Expected Shortfall (ES), respectively RORC or RORAC based on ES. In a pragmatic approach we combine and correlate a stock market model with geometric brownian motions with a two-factor Cox-Ingersoll-Ross (CIR-2) model for the interest rates/bonds. We use recent results from the theory of risk capital allocation, performance measurement and Swarm Intelligence for optimization. Examples for German market data as well as an analysis of the scalability of the solution to assure fast run-times on clusters of computers for large real-life portfolios are given.

**JEL:** G11, G31

**Keywords:** Cox-Ingersoll-Ross model; Expected Shortfall; Portfolio optimization; RORAC; Swarm Intelligence; Parallel computing

## 1 Introduction

In the first part of this paper we introduce a general concept of portfolio optimization with respect to the risk measure Expected Shortfall (ES) and also (but not at the same time) with respect to the performance measures RORC and RORAC (Return On Risk/Risk-Adjusted Capital) due to ES. Later, a specific financial market model that enables optimization of portfolios consisting of bonds and stocks is proposed. This model combines and correlates a stock market model of geometric brownian motions with a two-factor Cox-Ingersoll-Ross (CIR-2) model for the interest rates, respectively bonds. Numeric optimization is done using recent results from the theory of risk capital allocation, performance measurement and (Particle) Swarm Intelligence (SI). Explicit formulas and methods for the model and algorithms like Gradient Search (GS) and SI optimization are provided. Examples of optimized portfolios for German market data are given. Furthermore, we analyze the scalability of the solution (with respect to multi-processor machines, clusters or Global Grid environments) to assure fast run-times for large real-life portfolios.

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The idea of portfolio optimization with respect to modern risk and performance measures like ES, RORC or RORAC and also taking correlations between interest rates and stocks using an enhanced interest rate model into account seems to be new. Fast developing computer technology enables to solve optimization problems numerically even for complex market models and big portfolios.

Our approach is pragmatic in the sense that some of the theoretical problems which can emerge are (although not considered in depth) solved by unorthodox methods that seem to work well with real-life portfolios. The focus of the paper lies on the presented methods and not on the testing of these methods. From an academic point of view, this might be unsatisfying, but the mentioned models have (separately and sometimes in unfortunately simplified forms) been considered in some of the biggest German insurance companies. Practitioners should keep in mind that the paper shows what is actually possible in optimizing portfolios, but not how good the presented methods are from the historical point of view or compared to other market models. Such questions are not part of this paper.

The paper is organized as follows. Section 2 introduces the considered risk and performance measures. Section 3 explains the general approach to the optimization problem. Furthermore, the Gradient Search, the Swarm Intelligence optimization method and the role of stochastic simulation in these approaches are described. After a general introduction to optimization methods, Section 4 introduces and discusses the proposed financial market model. Section 5 is dedicated to the stochastic simulation part, i.e. the generation of market scenarios. In Section 6, information on parameter estimation can be found. Section 7 gives a brief chronological overview of all steps concerning our proposed optimization methodology. In Section 8 we present first results for German market data. Section 9 shows the scalability of our solution. In Section 10 we conclude. Finally in Appendix A we give two results on the form of the derivatives of Value-at-Risk and Expected Shortfall expressions.

We introduce some notation. Let us define the *total payoff*

$$X = X(u) := \sum_{i=1}^n u_i X_i \quad (1)$$

of a *portfolio*

$$u = (u_i)_{1 \leq i \leq n} \in \mathbb{R}^n \quad (2)$$

which represents  $n \in \mathbb{N}$  different *payoffs*  $X_i$  ( $1 \leq i \leq n$ ) with weights  $u_i \in \mathbb{R}$ . The  $X_i$  are assumed to be one-dimensional real-valued random variables. For example, each  $X_i$  can be considered as the difference of the price of asset  $i$  between the present time 0 and a future time horizon  $t > 0$ . Then the portfolio  $u$  of the different assets has a difference in value from time 0 to  $t$  which is exactly  $X(u)$ , and  $X > 0$  ( $X < 0$ ) is an increase (decrease) of the portfolio value from time 0 to  $t$  and therefore a win (loss) for the portfolio holder. We call  $B = (X_1, \dots, X_n)$  a *portfolio base* (cf. Fischer, 2003) as any considered portfolio will be described with (2) and (1). As random variables, the components of  $B$  *do not* have to be linearly independent.

## 2 Risk and performance measures

### 2.1 General definition

A *risk measure*  $\rho$  is usually defined as a mapping from a set of random variables (i.e. payoffs)  $\mathcal{X}$  to the real numbers, that means

$$\begin{aligned} \rho : \mathcal{X} &\longrightarrow \mathbb{R} \\ X &\longmapsto \rho(X). \end{aligned} \tag{3}$$

The amount  $\rho(X)$  is commonly interpreted as the minimum cash such that the “risk” of  $X$  is “acceptable” to the holder of the payoff or portfolio whenever he/she has the additional amount  $\rho(X)$  stored as *risk capital* (cf. Artzner et al., 1999).

Working with a portfolio base  $B = (X_1, \dots, X_n)$ , a risk measure  $\rho$  on the payoffs  $\mathcal{X}$  implies a risk measure  $\rho_B$  on the portfolios  $u$  out of  $\mathbb{R}^n$  for which we have  $X(u) \in \mathcal{X}$ . In particular, if  $X(\mathbb{R}) = \mathcal{X}$  we can define

$$\begin{aligned} \rho_B : \mathbb{R}^n &\longrightarrow \mathbb{R} \\ u &\longmapsto \rho(X(u)). \end{aligned} \tag{4}$$

We also write  $\rho(u)$  for  $\rho_B(u)$ . Based on the context, no confusion arises.

A *performance measure* can also be represented by functions as considered in (3) and (4). In contrast to risk measures, performance measures are usually intended to describe ratios like the relation of the expected return to the risk capital or invested risk-adjusted capital. However, the concrete interpretation of such measures is postponed until we look at concrete examples, namely RORC and RORAC (cf. section 2.3).

### 2.2 Expected Shortfall

For  $0 < \alpha < 1$ , we define *Value-at-Risk* as

$$\text{VaR}_\alpha(X) := -\inf\{x : P(X \leq x) \geq \alpha\}. \tag{5}$$

Hence, VaR is a negative  $\alpha$ -quantile of the distribution of the random variable  $X$ . *Expected Shortfall (ES)* is defined by

$$\text{ES}_\alpha(X) := -\mathbf{E}[X | X \leq -\text{VaR}_\alpha(X)]. \tag{6}$$

The meanings of these risk measures are obvious:  $-\text{VaR}$  is a threshold which is fallen short of in  $\alpha \cdot 100\%$  of all cases,  $-\text{ES}$  is the expectation (i.e. the mean) of the losses under the condition that this threshold has already been fallen short of. The change of the sign is a matter of interpretation – to neutralize losses (negative wins), risk capital has to be positive.

There are good reasons to only consider the ES risk measure. Ongoing from the widely known Value-at-Risk methodology, ES is easy to understand and always more conservative than VaR. Furthermore, ES is in most relevant cases a coherent risk measure (cf. Acerbi and Tasche, 2002) and features (when differentiable) explicit expressions for partial derivatives which is crucial in the context of risk capital allocation problems, but also for portfolio optimization which will soon become clear. We cite a risk management expert from the German Federal Reserve (*Deutsche Bundesbank*): “In my opinion, ES is still the *best* risk measure of all.”

## 2.3 RORC and RORAC

We define a performance measure

$$\varphi(X) := \frac{\mathbf{E}[X]}{\rho(X)}. \quad (7)$$

$\varphi$  is called the RORC, i.e. the *Return On Risk Capital*. In contrast to a risk measure, this performance measure does not care about the absolute value of the risk capital, but of its proportion to the mean return which is gained on it. For  $\rho = \text{ES}_\alpha$ , i.e.

$$\varphi_\alpha(X) := \frac{\mathbf{E}[X]}{\text{ES}_\alpha(X)}, \quad (8)$$

we talk of the ES-RORC. Some authors (cf. Tasche, 1999) call (7) the RORAC. We think that the “Return On Risk-Adjusted Capital” should be defined as in (10).

Working on RORC optimization, one might face the problem that the optimal portfolio (although the portfolio value is constant) implies a huge amount of risk capital together with a huge expected return. However, practical reasons might imply an upper bound for the risk capital. So we need a constraint  $\rho(u) \leq \rho_{\max}$ , i.e. in case of the ES-RORC

$$\text{ES}_\alpha(u) \leq \text{ES}_{\max}, \quad (9)$$

might be imposed.

The performance measure

$$\psi(X) := \frac{\mathbf{E}[X]}{V + \rho(X)} \quad (10)$$

is called the RORAC, i.e. the *Return On Risk-Adjusted Capital*. Indeed,  $\psi$  measures the mean (or expected) return per unit engaged capital, since  $V + \rho$  is the value of the invested capital plus the costs of risk. Hence, in contrast to RORC (7), RORAC considers not only the risk capital but the risk-adjusted investment capital and therefore seems to be a more sophisticated performance measure. For  $\rho = \text{ES}_\alpha$ , i.e.

$$\psi_\alpha(X) := \frac{\mathbf{E}[X]}{V + \text{ES}_\alpha(X)}, \quad (11)$$

we talk of the ES-RORAC.

As in the case of ES-RORC, the additional constraint (9) might be imposed in the case of ES-RORAC optimization.

## 3 Portfolio optimization

### 3.1 The problem

In this section we explain from a general point of view how to optimize a portfolio with respect to ES or with respect to the performance measures ES-RORC or respectively ES-RORAC.

We assume a fixed budget/portfolio value of  $V(u)$  which must be fully invested at the present time  $t = 0$ . Otherwise, the considered problems become trivial or unsolvable as they are usually linear in the portfolio value.

Let us assume that a portfolio  $u \in \mathbb{R}^n$  is given. Furthermore, this portfolio has to be optimized with respect to a risk or performance measure  $\rho$  on  $\mathbb{R}^n$ . For convenience, we assume that the risk  $\rho$  has to be minimized. As mentioned, exactly the amount  $V$  has to be invested in the market at time 0. If  $V_i$  is the price of asset  $i$  at time 0, this implies the following constraint for the portfolios  $u$  which must be satisfied:

$$V = \sum_{i=1}^n u_i V_i. \quad (12)$$

The solution of the optimization problem is given by a portfolio  $u^* \in \mathbb{R}^n$ , such that  $\rho(u^*)$  is minimal (on  $\mathbb{R}^n$ ) and the constraint (12) is fulfilled. Defining

$$u'_n := (V - \sum_{i < n} u_i V_i) / V_n, \quad (13)$$

and  $\rho'$  as

$$\rho'(u_1, \dots, u_{n-1}) := \rho(u_1, \dots, u_{n-1}, u'_n) \quad (14)$$

it follows from (12) that we can express the solution  $u^*$  by

$$(u_1^*, \dots, u_{n-1}^*) = \operatorname{argmin} \rho'(u_1, \dots, u_{n-1}), \quad (15)$$

together with  $u_n^* = (V - \sum_{i < n} u_i^* V_i) / V_n$ .

Working with real data, we discovered that portfolios which are extremal points due to the considered risk or performance measures can contain tremendous amounts of short-sold assets, i.e. the portfolio as a vector of real numbers contains huge negative components. For this reason we introduce a further constraint: For  $a, b \geq 0$  we require

$$a \frac{V}{V_i} \geq u_i \geq -b \frac{V}{V_i} \quad \text{for all } 1 \leq i \leq n. \quad (16)$$

For instance,  $b = 0$  implies portfolios which allow no short-selling. The values  $a = b = 1$  guarantee that the amount of capital or debts in no asset is bigger than the total value  $V$  of the portfolio.

As an optimization (e.g. for a 1-year horizon) can be driven daily or hourly, one could also think of *self-financing ES-*, *ES-RORC* or *ES-RORAC-optimal strategies* in this context.

### 3.2 Gradient Search

Let the risk measure (function)  $\rho$  be differentiable on  $\mathbb{R}^n$ . From standard analysis we obtain for  $1 \leq i \leq n - 1$

$$\frac{\partial \rho'}{\partial u_i}(u_1, \dots, u_{n-1}) = \frac{\partial \rho}{\partial u_i}(u_1, \dots, u_{n-1}, u'_n) - \frac{V_i}{V_n} \frac{\partial \rho}{\partial u'_n}(u_1, \dots, u_{n-1}, u'_n). \quad (17)$$

Using the partial derivatives (17), one can start looking for the (local) extreme points of  $\rho'$  in  $\mathbb{R}^{n-1}$  by applying Gradient Search (GS) methods. This might be a comfortable approach to solve the optimization problem (15) as long as the considered measures have sufficient differentiability properties. However, the proof of such differentiability properties can be rather difficult (cf. Appendix A or Tasche, 1999). This is one reason for our proposal of

```

maxits=bigNumber
its=0
while its < maxits      # fixed number of iterations
                        # or grid-based search
  ptf=RandomPortfolio # satisfying constraints
  loop
    if Rho(ptf)<Rho(bestPtf)      # small Rho wanted
      bestPtf=ptf
    end
    grd=gradient(ptf)
    ptfnew=ptf-grd*stepsize      # if constraints, adapt this step
  until Rho(ptfnew)>Rho(ptf)
  its+=1
end

```

Figure 1: Slow and simple Ruby-Pseudocode for portfolio optimization using brute-force gradient search. Clever varying choice of epsilon to calculate the gradient and the step-size can give further speed-up. Instead of choosing random portfolios, one can use a grid-based search.

Swarm Intelligence optimization methods (see Subsection 3.3).

Outline of a gradient minimum-search (Figure 1):

1. Evaluate the gradient for the current portfolio.
2. If the gradient is zero, exit. We have found a (local or global) minimum.
3. Follow the negative gradient (negative slope) of the current portfolio one small step. Modify the portfolio to satisfy the constraints. Then continue with step 1.

On a one-processor machine this gradient algorithm has to be started many times with different portfolios (“brute-force”), as one might be stuck in a local minimum. This takes a long time for real-life portfolios.

The following three paragraphs derive the respective partial derivatives (17) for Expected Shortfall, ES-RORC and ES-RORAC.

### Expected Shortfall

Assuming sufficient differentiability properties, results of Tasche (1999) show that

$$\frac{\partial \text{ES}_\alpha}{\partial u_i}(u_1, \dots, u_n) = -\mathbf{E}[X_i | X \leq -\text{VaR}_\alpha(X)]. \quad (18)$$

We refer to Tasche (1999) but also Lemma A.1 and A.2 in Appendix A for further information on the differentiation of the ES. For  $\rho(u) = \text{ES}_\alpha(X(u))$ , equation (17) can be written as

$$\begin{aligned} \frac{\partial \text{ES}'_\alpha}{\partial u_i}(u_1, \dots, u_{n-1}) &= -\mathbf{E}[X_i | X \leq -\text{VaR}_\alpha(X)] \\ &\quad + \frac{V_i}{V_n} \mathbf{E}[X_n | X \leq -\text{VaR}_\alpha(X)]. \end{aligned} \quad (19)$$

In fact, this simple expression of expectations is very suitable for numerical computations by Monte-Carlo methods.

### RORC

The partial derivatives (17) of the ES-RORC under the portfolio constraint (12) are obtained using standard rules of differentiation:

$$\begin{aligned} \frac{\partial \varphi'_\alpha}{\partial u_i} &= -\mathbf{E}[X] \cdot \frac{\mathbf{E}[X_i|X \leq -\text{VaR}_\alpha(X)]}{\text{ES}_\alpha(X)^2} + \frac{\mathbf{E}[X_i]}{\text{ES}_\alpha(X)} \\ &\quad - \frac{V_i}{V_n} \left( -\mathbf{E}[X] \cdot \frac{\mathbf{E}[X_n|X \leq -\text{VaR}_\alpha(X)]}{\text{ES}_\alpha(X)^2} + \frac{\mathbf{E}[X_n]}{\text{ES}_\alpha(X)} \right). \end{aligned} \quad (20)$$

As in the case of (19), we see from the definition (6) of ES that (20) is a relatively simple expression of expectations.

### RORAC

As  $V$  is a constant, the partial derivatives (17) of the ES-RORAC under the portfolio constraint (12) are similar to those of RORC (20):

$$\begin{aligned} \frac{\partial \psi'_\alpha}{\partial u_i} &= -\mathbf{E}[X] \cdot \frac{\mathbf{E}[X_i|X \leq -\text{VaR}_\alpha(X)]}{(V + \text{ES}_\alpha(X))^2} + \frac{\mathbf{E}[X_i]}{V + \text{ES}_\alpha(X)} \\ &\quad - \frac{V_i}{V_n} \left( -\mathbf{E}[X] \cdot \frac{\mathbf{E}[X_n|X \leq -\text{VaR}_\alpha(X)]}{(V + \text{ES}_\alpha(X))^2} + \frac{\mathbf{E}[X_n]}{V + \text{ES}_\alpha(X)} \right). \end{aligned} \quad (21)$$

## 3.3 Swarm Intelligence

Swarm Intelligence (SI) is a property of a system where the collective behaviours of (un-sophisticated) agents interacting locally with their environment cause coherent functional global patterns to emerge. SI provides a basis with which it is possible to explore collective (or distributed) problem solving without centralized control or the provision of a global model (cf. Kennedy et al., 2001).

The three underlying principles of SI are: evaluate, compare and imitate. Living organisms can learn by evaluating stimuli and rate them as positive or negative. In our case this is the metric (i.e. risk or performance measure) we want to minimize/maximize. As practiced in the Adaptive Culture Model (cf. Shibanaï, Yasuno and Ishiguro, 2001) and in real life, people compare themselves to others and imitate only those neighbours that are superior to themselves. Imitation is central to human sociality and important for the acquisition and maintenance of mental abilities (cf. Kennedy et al., 2001). SI offers a tradeoff between individual and group learning.

We give a brief outline of the algorithm (cf. Kennedy et al., 2001; Kennedy and Eberhart, 1995) and use standard notation. Let  $y_i$  be the position of particle  $i$ . In our case the position represents a specific portfolio ( $y_i \in \mathbb{R}^n$ ). The change of portfolio is called  $v$ .  $v$  traditionally stands for velocity. Each clockstep  $t$  particles move from one stop to another by  $y_i(t) = y_i(t-1) + v_i(t)$  and sample the search space by modifying the velocity term. The direction of movement is a function of the current position ( $y_i$ ), velocity ( $v_i$ ), the location of the individual's previous best success ( $p_i$ ), and the best position found by any member of the neighborhood ( $p_g$ ):

$$y_i(t) = f(y_i(t-1), v_i(t-1), p_i, p_g). \quad (22)$$

One possible implementation is

$$v_i(t) = v_i(t-1) + n_1(p_i - y_i(t-1)) + n_2(p_g - y_i(t-1)) \quad (23)$$

with

$$y_i(t) = y_i(t-1) + v_i(t). \quad (24)$$

The  $n$  variables are random variables defined by an upper limit, so that the particles cycle around the two best bets:  $p_i$  and  $p_g$ . The random numbers ( $n_1$  and  $n_2$ ) are updated in every iteration. With real-life data the velocity  $v$  very quickly becomes too large and one has to set limits.

```

ys=generateInitialPortfolios      #satisfying the constraint
p=ys
loop
  # best portfolios's fitness so far
  ys.each_with_index{ |y,i|
    p[i]=y, if Rho(y)<Rho(p[i]) # small Rho wanted
  }

  i=rand size                      # arbitrary Rho
  g=i
  for j=indexes of neighbors
    g=j if Rho(p[j])<Rho(p[g]) # g: index of best performer
                                # in the neighborhood
  end
  #assuming delta t=1
  v[i]=[i]+n1*(p[i]-ys[i])+n2*(p[g]-ys[i])
  v[i]=Vmax if v_id>Vmax
  v[i]=Vmin if v_id<Vmin
  ys[i]=ys[i]+v[i]

  fixPortfolio                     # constraint
  if loopCount mod 10000==0       # big number here
    removeWorstPortfolios        # remove 10% worst portfolios
    injectNewPortfolios          # inject 10% new portfolios
  end
until some criterion

```

Figure 2: Extended and slow but simple Ruby-Pseudocode for portfolio optimization using swarm particles based on Kennedy et al. (2001). This basic algorithm is implemented more efficiently.

As the present value of a portfolio has to remain constant, two minor modifications in the choice of  $v$  are required.

In simulation studies on typical portfolios it proves successful to inject about 10% of new particles with random speeds and locations from time to time and to remove the 10% worst performing particles. The exact population size is an open research problem with experts having different opinions. A rule of thumb is to keep the population size small, but to rely on a high number of iterations. As this can take a long time for higher dimensional problems, parallel solutions are an easy way out of the dilemma, following Kent Thompson's (co-inventor of Unix) famous quote: "When in doubt, use brute force".



### 3.4 How stochastic simulation fits in

Independent from the question whether GS or SI methods are used to solve the portfolio optimization problem, it is clear that a way must be found to determine the distributions of the considered payoff functions  $X_i$  (cf. Section 1, equation (1)) as finally in any optimization routine the risk or performance measures (6), (8) and (11) must be computed. The  $X_i$  were interpreted as wins or losses due to the  $i$ -th asset in the market where we assumed to have  $n$  numbered assets. Let us assume that the prices of these assets at some time  $t \in \mathbb{R}_0^+$  are given by random variables

$$V_1(t), V_2(t), \dots, V_n(t). \quad (25)$$

We assume to have constants for  $t = 0$ , which is the present time. In particular, we used the notation

$$V_i = V_i(0) \quad (26)$$

and

$$X_i = V_i(t) - V_i(0) \quad (27)$$

until now. The variable  $t > 0$  is the considered time horizon for which risk management is performed.

It is clear that many of our thoughts so far, especially the functions ES, ES-RORC, ES-RORAC, but also their derivatives, crucially depend on the model for future prices  $V_i(t)$ . The particular stochastic model we use is introduced in Section 4.

Once the model for the price processes  $V_i$  is chosen and a way to get possible parameters is found, one can theoretically compute the risk and performance measures (6), (8) and (11) and their partial derivatives under budget constraint (19), (20) and (21). However, one often encounters models (also in our case) where it is not possible (or quite difficult) to compute these values directly. The more realistic assumption is that one succeeds in doing a stochastic simulation of the model which computes  $m \in \mathbb{N}$  (e.g.  $m = 10^3$ ) market scenarios, i.e. finally one has for each  $i$  the numerical realizations (in increasing order)

$$x_i^1, x_i^2, \dots, x_i^m \quad (28)$$

of the random variable  $X_i$  defined by (27). The realizations (also in increasing order)

$$x^1, x^2, \dots, x^m \quad (29)$$

for any  $X = X(u)$  follow immediately.

Having these realizations, estimates for the stochastic expressions in the functions mentioned above can be used. In particular, we compute estimates using the “empirical” distribution given by the simulation output, e.g.

$$\widehat{\mathbf{E}}[X] = \sum_{j=1}^m x^j / m, \quad (30)$$

$$\widehat{\text{VaR}}_\alpha(X) = -x^{\lfloor \alpha m \rfloor} \quad (31)$$

or

$$\widehat{\text{ES}}_\alpha[X] = \sum_{x_j \leq -\widehat{\text{VaR}}_\alpha(X)} x^j / m, \quad (32)$$

where  $[r]$  denotes the integer part of any real number  $r$ . Of course, one can use other (perhaps more sophisticated) estimators. Nonetheless, replacing all stochastic expressions in (6), (8), (11), (19), (20) and (21) as suggested by (30) to (32), one obtains approximations of the respective measures and their gradients which are easy to implement in any suitable programming language.

Gradient Search methods or Swarm Intelligence optimization methods can now be executed using the obtained approximations.

## 4 The proposed market model

Until now, the presented theory has not been fixed to a particular financial market model and was intended to give a general introduction to the portfolio optimization problem. In the following sections we apply the above ideas to a concrete model and data setup.

We model stocks and non-defaultable bonds. All stochastics evolves from a  $(d + 2)$ -dimensional brownian motion  $(B_i)_{i=1,\dots,d+2}$ , where the first two components drive the dynamics of the two-factor interest rate model for the bonds and the last  $d$  drive the dynamics of  $d$  stocks. The brownian motions are correlated by a covariance matrix  $\Sigma$ .

### 4.1 Interest rates and bonds

We use

$$e^{R(t,\tau)\cdot\tau} = \frac{1}{p(t,\tau)} \quad (33)$$

as the defining equation of the relation between the price  $p(t,\tau)$  of a *zero-coupon bond* with maturity  $\tau$  at time  $t$ , i.e. the price at time  $t$  of the guaranteed payoff 1 at time  $t + \tau$ , and the corresponding *spot (interest) rate*  $R(t,\tau)$ . Hence,  $R(t,\tau)$  is the at  $t$  guaranteed continuous interest rate during the time interval  $[t, t + \tau]$ . For future points of time ( $t > 0$ ),  $p(t,\tau)$ , respectively  $R(t,\tau)$ , are assumed to be random variables. We now turn to the considered interest rate model of Chen and Scott (1992) with two stochastic factors.

The model is usually called Cox-Ingersoll-Ross-2 (CIR-2) as it relies heavily on the work of Cox, Ingersoll and Ross (1985) which is a so-called short rate model with only one (economically interpretable) stochastic factor (modelled by a square-root process). However, the authors also formulated the main ideas for a theory with multiple stochastic factors. In our description of the model, we closely follow Fischer, May and Walther (2003), which also includes comments on the model choice which we want to adopt for our purposes (see also Subsection 4.3).

The concrete model setup is given by the two stochastic factors  $x = (x_1, x_2)$  fulfilling the stochastic differential equations

$$dx_i = (b_i - a_i \cdot x_i)dt + \sigma_i \sqrt{x_i} dB_i \quad (i = 1, 2) \quad (34)$$

where  $b_i$ ,  $a_i$  and  $\sigma_i$  are positive constants. One has  $x_i > 0$  if  $2b_i > \sigma_i^2$ .  $B_i(t)$  is the  $i$ -th brownian motion at time  $t$ ,  $B_1$  and  $B_2$  are *independent* (not correlated). Equation (34) defines a so-called *mean reversion process*. The parameter  $a$  is called the *strength* of the mean reversion and  $b/a$  the *mean reversion level*, i.e. the long-term mean of the process  $x_i$ .

The implied spot interest rate at time  $t$  for a maturity  $\tau$  is

$$R(t, \tau) = \sum_{i=1}^2 \left( -\frac{\log A_i(\tau)}{\tau} + \frac{B_i(\tau)}{\tau} x_i(t) \right), \quad (35)$$

the implied zero-coupon bond price at time  $t$  for the maturity  $\tau$

$$p(t, \tau, x(t)) = \prod_{i=1}^2 A_i(\tau) e^{-B_{d+i}(\tau) x_i(t)}. \quad (36)$$

The respective functions  $A_i$  and  $B_i$  are given by

$$A_i(\tau) = \left[ \frac{2h_i e^{(a_i + \lambda_i + h_i)\tau/2}}{2h_i + (a_i + \lambda_i + h_i)(e^{\tau h_i} - 1)} \right]^{2b_i/\sigma_i^2} \quad (37)$$

and

$$B_i(\tau) = \left[ \frac{2(e^{\tau h_i} - 1)}{2h_i + (a_i + \lambda_i + h_i)(e^{\tau h_i} - 1)} \right], \quad (38)$$

with

$$h_i = \sqrt{(a_i + \lambda_i)^2 + 2\sigma_i^2}. \quad (39)$$

The parameter  $\lambda_i$  is concerning the change of measure (physical to martingale measure) and can also be estimated from historical interest rates. In the one-factor case, a particular function of  $\lambda$  is interpreted as the so-called market price of risk (Cox, Ingersoll and Ross, 1985; see also Fischer, May and Walther, 2002). For more than one factor, an economic interpretation is not possible (or at least not obvious).

It is clear that the price of any coupon bond can be computed as the sum of the prices of the respective set of zero-coupon bonds.

## 4.2 Stocks

The  $d$  stocks of the considered financial market are modelled by geometric brownian motions, i.e. price processes  $S_j$  ( $j = 1, \dots, d$ ) with

$$S_j(t) = S_j(0) e^{\mu_j t + \sigma_j B_{j+2}(t)}, \quad (40)$$

where  $\mu_j \in \mathbb{R}$  is the drift and  $\sigma_j \in \mathbb{R}^+$  the diffusion coefficient of the brownian motion in the exponent, i.e. the price process has the trend  $\mu_j + \sigma_j^2/2$ . In terms of stochastic differential equations (SDE) we have

$$d \ln S_j = \mu_j dt + \sigma_j dB_{j+2}. \quad (41)$$

$B_i(t)$  is the  $i$ -th brownian motion at time  $t$ .

## 4.3 Comments

The model choice is based on our experience with practitioners. We know that at least in three major German life insurance companies one-factor Cox-Ingersoll-Ross models together with geometric brownian motions have been considered in the context of Asset Liability Management. CIR-1 is used to model the debt securities market and interest rates whereas the geometric brownian motions (in reminiscence of the Black-Scholes model) are used to

model “stocks”. The mentioned insurance companies are interested in multi-factor models although not yet using them. The combination and correlation of the models as proposed in this paper seems to be new.

For insurance companies (known to be conservative), an important aspect of such models is the acceptance by the scientific public. This enlightens the decision for standard models like geometric brownian motions or the CIR-1 model.

As mentioned in Fischer, May and Walther (2003), for instance the Vasicek-2 (Gaussian) model behaves in some way better than CIR-2 (concerning parameter estimation or the values of the likelihood function; see also Babbs and Nowman, 1998). Nonetheless, insurance companies seem to prefer CIR, as under the respective parameter constraints CIR assures positive interest rates. Facing a possible deflation in the Eurozone (especially in Germany), one might want to reconsider this philosophy.

From the academic point of view it is clear that alternative models like the Vasicek model should also be examined with respect to the optimization problem. However, for several reasons which will become clear later we recommend to stay inside the class of so-called affine term structure models.

Another more theoretical problem is whether the probably for optimization purposes used derivatives (19), (20) and (21) really exist. Depending on the considered model, this might not be trivial, see also Appendix A for some comments on the differentiability problem. We have not proven the existence of the derivatives for the proposed model (in this case, the Vasicek model may be easier to handle, too). However, for our purposes, this unsolved theoretical problem (which relies on the used model) is no drawback as our optimization routines are subject to “back-testing” by the SI methods. Nonetheless, the GS methods work very well in the search of local extrema.

The problems coming in line with differentiation of quantile expressions could be avoided by using risk measures which have more suitable differentiability properties as e.g. the risk measures depending on one-sided moments which are proposed in Fischer (2003).

The authors admit that the proposed model has not been examined for absence of arbitrage. This is postponed to further research. Actually, the model is used like an econometric framework. In this sense, the philosophy of our approach is pragmatic.

## 5 Market scenario generation

As described in Subsection 3.4, we carry out a stochastic simulation to obtain an “empirical” distribution of the considered random payoffs.

A simulation requires discretization. We consider points of time  $t_m$  ( $m \in \mathbb{N}_0$ ). The width of the time step is the constant  $\Delta$ , e.g. one day, one month etc., i.e.  $t_{m+1} = t_m + \Delta$  and  $t_0 = 0$ . Increments

$$\delta B_{i,m} := B_i(t_m) - B_i(t_{m-1}) \quad (42)$$

of the  $(d+2)$  brownian motions have to be simulated. For fixed  $m$ , the  $\delta B_{i,m}$  are correlated by a common covariance matrix  $\Delta\Sigma$ . For fixed  $i$ , the increments  $\delta B_{i,m}$  are independent normally distributed random variables with variance  $\Delta$  and expectation 0.

Hence, all discretized dynamics are driven by a series of standard normally distributed

random variables  $N_{i,m}$  ( $i = 1, \dots, d+2; m \in \mathbb{N}$ ), where for each  $m$  the random variables  $(N_{i,m})_{i=1, \dots, d+2}$  are correlated by the covariance matrix  $\Sigma$  which will later be estimated from real data.

### 5.1 Simulation of correlated normal random variables

Simulation of i.i.d. normal random variables is standard. Let us consider the Cholesky decomposition

$$\Sigma = CC^t \quad (43)$$

of the covariance matrix  $\Sigma$ . If  $Z = (Z_i)_{i=1, \dots, d+2}$  are  $d+2$  i.i.d. normal random variables, then

$$(N_i)_{i=1, \dots, d+2} = N = C \cdot Z \quad (44)$$

contains  $d+2$  normally distributed random variables with covariances  $\Sigma$ .

### 5.2 Interest rates and bonds

From (34) an Euler-approximation gives the recursion

$$x_{i,m} = x_{i,m-1} + (b_i - a_i \cdot x_{i,m-1})\Delta + \sigma_i \sqrt{x_{i,m-1}} \sqrt{\Delta} N_{i,m} \quad (i = 1, 2) \quad (45)$$

where the  $N_{i,m}$  (fixed  $i$  or alternatively fixed  $m$ ) are i.i.d.  $N(0, 1)$  (see also Fischer, May and Walther, 2003). For a general introduction into the numerics of stochastic differential equations we refer to Kloeden and Platen (1992).

Plugging the computed values into (35), resp. (36), returns the desired interest rates, resp. bond prices.

### 5.3 Stocks

From (41) we get the Euler-approximation

$$\ln S_{j,m} - \ln S_{j,m-1} = \mu_j \cdot \Delta + \sigma_j \cdot \sqrt{\Delta} \cdot N_{j+2,m} \quad (1 \leq j \leq d) \quad (46)$$

where the  $N_{j+2,m}$  (fixed  $j$ ) are i.i.d.  $N(0, 1)$ . This implies for  $M \in \mathbb{N}$

$$S_{j,M} = S_{j,0} \exp \left( \mu_j M \Delta + \sigma_j \sum_{m=0}^M \sqrt{\Delta} N_{j+2,m} \right). \quad (47)$$

## 6 Estimation of parameters

### 6.1 Interest rates and bonds

The estimation of the parameters of the CIR-2 model and detailed description of the used methods are subject of several existing articles, e.g. Chen and Scott (1993), Duan and Simonato (1999), Bolder (2001), Beletsky and Szimayer (2002) and Fischer, May and Walther (2003). The problem is not trivial. The most efficient method seems to be maximum-likelihood estimation with Kalman-filtering. In particular, we used the machinery as explained in Fischer, May and Walther (2003). The interested reader can find further information in this paper and the references therein.

A comment on the data: We use the historical yield structure of the German debt securities market (monthly, taken at the end of each month). The values for spot rates with maturities  $\tau > 0$  up to 28 years can be computed via a parametric presentation of yield curves (the so-called Svensson-method; cf. Svensson, 1994, and Schich, 1997) for which the historical parameters can be taken from the homepage of the German Federal Reserve (*Deutsche Bundesbank*; <http://www.bundesbank.de>). The implied Bundesbank values  $R'$  are estimates of *discrete* interest rates on notional zero-coupon bonds based on German Federal bonds and treasuries (cf. Schich, 1997) and have to be converted into continuous interest rates by  $R = \ln(1 + R')$ .

## 6.2 Stocks

Given the market data  $S_{j,m}$  ( $j = 1, \dots, d$ ;  $m = -M, \dots, 0$ ; time step =  $\Delta$ ;  $t = 0$  is the present), discretization (46) is used to compute the estimators

$$\hat{\mu}_j = \frac{1}{M\Delta} \ln(S_{j,0}/S_{j,-M}) \quad (48)$$

and

$$\hat{\sigma}_j = \sqrt{\frac{1}{M\Delta} \sum_{i=1-M}^0 (\ln(S_{j,m}/S_{j,m-1}) - \Delta\hat{\mu}_j)^2} \quad (49)$$

for the parameters of the stock price dynamics.

## 6.3 The covariance matrix

After having plugged in historical data, solving equations (45) and (46) for the values  $N_{i,m}$  gives us a time series  $(N_{i,m})$  ( $i = 1, \dots, d+2$ ;  $m = 1-M, \dots, 0$ ) of hypothetical historical realizations of the normal random variables (44). Now, the values

$$\hat{\Sigma}_{i,j} = \frac{1}{M} \sum_{m=1-M}^0 N_{i,m} N_{j,m} \quad (50)$$

can be used as estimates for the entries of the covariance matrix  $\Sigma$ . However, there is still something missing since we can not get the historical realizations  $x_{i,m}$  of the stochastic factors of the interest rate model (45) directly from the market. Instead, we use the affine term structure (35) to derive them from the interest data distributed by the German Federal Reserve (cf. Subsection 6.1). One has

$$\underbrace{\begin{pmatrix} R(t, \tau_1) \\ R(t, \tau_2) \end{pmatrix}}_{R_t} = \underbrace{\begin{pmatrix} -\frac{\log A_1(\tau_1)}{\tau_1} & -\frac{\log A_2(\tau_1)}{\tau_1} \\ -\frac{\log A_1(\tau_2)}{\tau_2} & -\frac{\log A_2(\tau_2)}{\tau_2} \end{pmatrix}}_{M_A} + \underbrace{\begin{pmatrix} \frac{B_1(\tau_1)}{\tau_1} & \frac{B_2(\tau_1)}{\tau_1} \\ \frac{B_1(\tau_2)}{\tau_2} & \frac{B_2(\tau_2)}{\tau_2} \end{pmatrix}}_{M_B} \cdot \underbrace{\begin{pmatrix} x_1(t) \\ x_2(t) \end{pmatrix}}_{x(t)} \quad (51)$$

$$R_t = M_A + M_B \cdot x(t). \quad (52)$$

Hence, we obtain by

$$x(t) = M_B^{-1}(R_t - M_A) \quad (53)$$

a time series  $x_{i,m}$  ( $i = 1, 2$ ;  $m = -M, \dots, 0$ ) by inserting the time series of the respective spot rates into (53). Slightly different from Fischer, May and Walther (2003), our suggestion is

$$\tau_1 = 0.5 \text{ years}, \quad \tau_2 = 10.0 \text{ years}. \quad (54)$$

Equation (53) also returns the starting values  $x(0) = (x_1(0), x_2(0))$  for the simulation of the factors  $x_1$  and  $x_2$ . The computation of the values  $x(0)$  implies a (mathematically) continuous continuation of the history of the spot rates  $R(\cdot, \tau_1)$  and  $R(\cdot, \tau_2)$  by the CIR-2 model. For other maturities than  $\tau_1$  and  $\tau_2$  there might be jumps in the dynamics of the respective sport rate (cf. Fischer, May and Walther, 2003). A simulation study of the same authors showed that for realistic time horizons the starting values have significant influence on the means of the simulated interest rates. Hence, a proper calculation of starting values is important.

Having executed the explained procedure, one can compute the empirical covariance matrix  $\widehat{\Sigma}$  by (50). At this point, a further problem arises. The CIR-2 model works with *uncorrelated* brownian motions (cf. subsection 4.1). Nonetheless, the upper left  $2 \times 2$ -submatrix of  $\widehat{\Sigma}$ , which theoretically should be the two-dimensional identity, may differ from the theoretical values. To stay in the proposed model, one can adjust the estimate  $\widehat{\Sigma}$  by setting the upper left  $2 \times 2$ -submatrix to the identity matrix. Doing this, it is important to check whether the new matrix is still positively definite as we afterwards have to carry out the Cholesky decomposition. In cases where positive definiteness gets lost, one should choose a symmetric positively definite matrix close to the proposed matrix with the identity in the upper left corner.

The proposed technique for the computation of the covariance matrix and the starting values should be suitable for any stochastic interest rate model with an affine term structure as in (35) (e.g. Vasicek-2).

## 7 Chronological overview

### I. Estimation

- Get data.
- Estimate parameters of stock prices; (48) and (49).
- Estimate parameters of interest rate dynamics (cf. Subsection 4.1).
- Compute the historical time series  $x_{i,m}$  ( $m \leq 0$ ) by (53).
- Solve equations (46) and (45) for the historical  $N_{j,m}$  ( $m \leq 0$ ).
- Compute the covariance matrix  $\widehat{\Sigma}$ ; (50).
- Compute the Cholesky decomposition of  $\widehat{\Sigma}$ ; (43).

### II. Simulation (one loop)

- Simulate future i.i.d. normal random variables and plug them into (44) to get the simulated  $N_{i,m}$  ( $m > 0$ ).
- Plug the  $N_{i,m}$  into (46) and (45) to get the simulated scenarios of stock prices and interest rate model factors  $x_{i,m}$  ( $m > 0$ ).

- Plug the factors  $x_{i,m}$  into (35) or (36) to get spot rates or bond prices.
- Compute portfolio values, e.g. by (1).

### III. Evaluation

- Compute the risk and performance measures (6), (8) and (11) by the empirical distributions obtained by reiteration of step II; cf. (30) to (32).
- If necessary, compute the partial derivatives (19), (20) and (21).

### IV. Optimization

- Use a GS or SI method repeating step III for each new portfolio.

## 8 First results

### 8.1 Gradient Search vs. Swarm Intelligence

(Particle) Swarm Intelligence is a powerful tool to solve optimization problems in a fixed search-space. SI is computationally appealing as simple to implement and computationally robust with respect to local minima and maxima, provided enough iterations (generations) are performed. As an additional bonus, SI is inherently parallel and can be implemented in a massively parallel way (cf. Auslander et al., 1995, Fabiunke, 2002).

Gradient (Grid) Search methods like hill-climbing are superior to random-guessing algorithms like SI if the search-space is e.g. a sphere, but on highly multi-dimensional surfaces, the gradient method gets stuck too often in local extreme points and therefore becomes computationally expensive, as one has to start from many different starting points.

In higher-dimensional problems, SI seem fitter than GS methods. However, one has to be careful with such statements, as according to the No Free Lunch (NFL) theorem (cf. Wolpert and Macready, 1996), when performance is averaged over all possible search spaces, all search algorithms perform equally well.

Ultimately we decided to stick with SI algorithms, which seem to converge faster for large real-life size portfolios. Combining these evolutionary algorithms with a selected Gradient Search for selected good intermediate solutions provides further speed-up.

One caveat with all numerical solutions, without further assumptions about the search-space is that there is no guarantee that the optimal solution is found. In practice one monitors the rate of convergence and dedicates enough search-time. Looking at the number of idle PCs and workstations in the typical investment bank or insurance company one can be on the save side and farm out the work in fractions of a second to a large number of processors or a dedicated cluster or Global Grid.

### 8.2 Examples

The general setup for our numeric examples is a time horizon of one month where the simulation takes 20 steps per month. The number of loops is 1000. We consider portfolios which have a present value of exactly 1000 EUR. We optimize using a local GS method and a combined GS-SI method. The second one is run with and without constraint  $b = 0$ ,



i.e. with and without shortselling in the portfolio (cf. (16)). The considered confidence level is 5%. Two types of portfolios are examined. The smaller one contains two bonds and two stocks, the bigger one 10 bonds and 10 stocks. In particular, we considered the following bonds and stocks (which are here listed in the same order as in the portfolio vectors):

- 2 zero-coupon bonds: Maturity 1 year and 10 years.  
2 “stocks”: Xetra DAX and Allianz
- 10 zero-coupon bonds: Maturity 1 year up to 10 years.  
10 stocks: Allianz, BASF, BMW, Bayer, Commerzbank, DaimlerChrysler, Deutsche Bank, Lufthansa, E.on, Hypovereinsbank

All stocks are elements of the Xetra DAX and had their IPO (Initial Public Offering) at least 10 years ago. Data was taken from <http://de.finance.yahoo.com>. The estimates are calculated from monthly data from May 2002 to April 2003. We obtain the model parameters listed in Table 1. The same time interval and discretization was taken for the estimation of the term structure model parameters (cf. Subsection 6.1). Maturities from 1 to 10 years were taken into consideration. Results are in Table 2.

The values of the (adjusted) covariance matrix in Table 3 confirm the use of correlations between the interest rate model factors and the stock market dynamics to obtain a more realistic combined model.

For each of our setups we computed the ES-, ES-RORC- and ES-RORAC-optimal portfolio. The mean, VaR, ES, ES-RORC and ES-RORAC for these portfolios are listed in the Tables 4-9 in Appendix B (and the portfolios themselves in the four assets case). The optimized portfolios are compared with “normed” portfolios where the same capital is invested in each of the four, respectively 20 assets. As expected, all optimized measures have been improved significantly (see also Figures 3 and 4) and the combined GS-SI method is superior to the local GS method starting at the normed portfolio. Local extreme points seem to exist in the most considered cases. A situation as in Table 6 where the ES of the ES-optimized portfolio is lower than (but close to) the ES of the RORC-optimized portfolio could be a symptom for the need of more (or finer) iterations.

Due to our pragmatic approach we did not invest any time in proofs for the existence or absence of global, respectively local extreme points in our model. Real financial companies are not interested in such questions, especially as portfolios are often optimized in small steps (and not a complete restructuring).

An interesting (and reasonable) model output is that the local GS results imply that bonds of longer maturities bear more financial risks. This can be seen in decreasing weights of bonds with higher maturities in the optimized portfolios (this is also true for the portfolios with 20 assets which are not listed in detail).

Massive short-selling and probable absence of global extreme points (e.g. in the RORAC case, cf. Table 4) motivate the use of constraint  $b = 0$  (no short-selling). Roughly speaking, the implication seems to be that optimized portfolios under the constraint contain almost no stocks. Optimization under the no-shortselling constraint seems to imply rather similar optimized portfolios for all measures (cf. Tables 6 and 9).

In summary, all obtained results seem to be reasonable from the economic point of view and confirm the proposed methods. We cannot really judge the impact of other models at this stage of our research. However, we guess that reasonable models (e.g. such using Vasicek-2) will imply results close to ours.

## 9 Parallel programming and scalability

### 9.1 Bulk synchronous parallel computing

Since 1944 von Neumann's model for sequential computing has been widely accepted, but there is no standard model for parallel computing. Most approaches nowadays are based on message-passing, but they are often inadequate, since the potential danger of deadlock, in which each possible activity is blocked, waiting on some other activity that is also blocked, increases dramatically with the complexity of software.

Furthermore, models based on message-passing, e.g. MPI (Message passing interface), do not easily allow performance prediction. The Bulk synchronous parallel computing model (BSP) however abstracts low-level program structure in favour of so-called supersteps. This allows easy debugging, removes the problem of deadlock and allows a reasoning of the correctness of the code nearly as easily as in sequential code.

#### BSP computers

A BSP computer consists of a set of processor/memory pairs, a global communication network and a mechanism for the efficient barrier synchronisation of the processors. In real life, this could be anything: a single/multi-processor PC, a cluster of workstations or a real parallel machine like the Cray T3D.

#### Supersteps

The fundamental idea of BSP is the notion of a superstep. In a superstep, computation and communication are decoupled. This avoids deadlock.

First the processes perform as many calculations as possible using their local data. If one processor needs data from another, communication starts only after all the computation has stopped. When communication is finished, barrier synchronisation is called and the next superstep begins.

### 9.2 Cost modelling and performance prediction

A cost model helps to guide the choice of programming algorithm.

The separation of communication from synchronisation and the inherent simplicity of the superstep structure make it relatively easy to find a suitable cost-model. The cost is expressed in terms of *steps* or *floating point operations (FLOps)* for each portion of the program. The cost parameters are the BSP parameters for the machine and parameters determined by the choice of algorithm and their implementation.

As a BSP program consists of a sequence of supersteps, the "cost" of an entire program is the sum of the contributions from its supersteps.

What are the key parameters that determine performance? Extensive research by the originators of the BSPlib showed that the following four key parameters are sufficient (cf. Hill and McColl, 1996):

- the number of processors,  $p$ ;
- processor speed,  $s$ , (number of steps per second);
- the cost  $l$  (steps), of achieving barrier synchronisation (which depends on network latency, which is a measurement of delay from one end of a network to another). Basically  $l$  is the cost of telling all processors to wait till all communication has been performed; and
- the cost  $g$  (steps per word), of delivering message data. This captures the interprocess communication speed.

Since the processor speed  $s$  is essentially a normalising factor, there are only three independent parameters:  $p$ ,  $l$  and  $g$ .

The cost of one superstep is

$$\max(w_i) + g \cdot \max(h_i) + l \tag{55}$$

where  $i$  ranges over processors ( $i = 1, \dots, p$ ),  $w_i$  is the time for the local computation in processor  $i$  and  $h_i$  is the number of incoming or outgoing messages per processor. The values of the parameters are determined by measurement using suitable benchmarks that mimic average computation and communication loads (cf. Hill, 1996).

The dependence on a specific platform enters the cost function only through the parameters  $p$ ,  $l$  and  $g$ .

We follow convention and count every floating point operation as 1.

The BSP approach offers a simple cost model. In general, cost-modeling applications give a rough ball-park figure of the cost on any parallel machine and configuration size. The role of profiling tools like `bsprof` aids simplistic pencil and paper cost modeling, and it effectively predicts the cost of an algorithm on any parallel machine (cf. Hill, Crumpton and Burgess, 1996).

### 9.3 Scalability

Programmers take the burden of writing parallel programs to increase speed and memory. The aim of every parallel algorithm designer is to write code that scales linearly, i.e. runs  $p$  times as fast on a  $p$ -processor machine. This clearly constitutes an upper bound, if the sequential algorithm is already optimal. Linear scalability is achieved by using good load-balancing, keeping all processors busy all the time and communication costs are minimized.

Data dependency can make optimal speed-up impossible. It determines parallel complexity, the minimum number of steps an algorithm would need to run on a PRAM-computer. This constitutes an upper bound on the maximal speed-up that can be achieved.

There are many different and more sophisticated layouts of parallel implementations possible. The right choice depends on the size of the portfolio and available hardware. For

the sake of simplicity in this article we have chosen the brute force approach.

Sketch of the scalability for a parallel brute-force GS: To avoid local extrema, one has to start many times from different grid-points:

1. Superstep: Broadcast the initial portfolio structure and search-areas, or only the portfolio structure and use random startpoints. Depending on the network architecture (reflected in the value of  $g$ ), one might use several supersteps and use e.g. a tree-shaped communication form. Asymptotic cost for a 1 phase broadcast:  $l + npg$ , where  $n$  is the size of the initial portfolio structure.
2. Superstep: Now work out gradient searches on all processors for a given time. E.g. every processor performs a set-number of searches. On average this will balance out. Asymptotic cost:  $1/p \times$  sequential time, as if  $p$  processors work out  $k/p$  searches,  $k$  searches are performed in total. The sequential time is the all dominating factor.
3. Superstep: Each processor sends its best grid point back to processor 1, which sorts them and gives the final result. Asymptotic cost:  $l + ng + p$ . The extra  $p$  arise from choosing the point with best fitness.

As the communication cost, sorting, etc. is negligible for any reasonable number of searches, this algorithm clearly scales linearly with the number of processors used.

Sketch of the scalability for SI:

Superstep: As in the 1-processor mode (see Figure 2), but now performed on all  $p$  processors. Every 1000 or 10000 iterations fit values are exchanged, then the next superstep starts.

Since the cost of data interchange is negligible compared to the cost of the iterations in each superstep, we have scalability as in the GS case.

One typical schoolbook error in this context is not to use a high quality random number generator, assuring independent random number streams on all processors (cf. Mascagni, Ceperley and Srinivasan, 1998 and 1999).

Large clusters as well as the rise of grid-computing requires analytic forecastig of run-times to chose the appropriate hardware for the task. There are many potential trade-offs (cf. Jarvis et al., 2002, 2003, and Roehrl, 1998): time versus money, etc. Our paper has shown that a pragmatic approach can take advantage of developments in computerscience to enable the exploration of new portfolio optimization techniques using parallel computing techniques.

## 10 Conclusion

The purpose of this paper is twofold. First, we describe in Section 1 to 3 a general methodology of ES-, ES-RORC- and ES-RORAC-optimization which seems to be suitable independently from the considered market model. Second, we propose a particular market model

which seems to be suitable to describe at the same time bonds and stocks as well as dependencies between them and which is used for our numerical examples. We thoroughly explain the proposed model and the respective simulation and optimization procedures. Concrete examples and a scalability analysis show the suitability and practicability of the methodology. Alternative models, model tests and the examination of some more theoretical questions have to be postponed to future research.

## A Derivatives of VaR and ES

This section derives expressions for the derivatives of Value-at-Risk and Expected Shortfall. Equation (18) is directly implied by Lemma A.2.

We consider a bivariate random variable  $(X, Y)$  with continuous density  $f(x, y)$  such that  $X + \epsilon Y$  has for any  $\epsilon \in \mathbb{R}$  a continuous density, too. Define  $\text{VaR}_\alpha(\epsilon)$  for  $0 < \alpha < 1$  as

$$\text{VaR}_\alpha(\epsilon) := \text{VaR}_\alpha(X + \epsilon Y). \quad (56)$$

Since  $X + \epsilon Y$  is a continuous random variable, the infimum in (5) is actually reached (i.e. is a minimum) and the respective probability is exactly  $\alpha$ .

**LEMMA A.1.** *Under certain strong assumptions on the density  $f$ , the Value-at-Risk  $\text{VaR}_\alpha(\epsilon)$  can be differentiable in  $\epsilon$  and*

$$\frac{\partial \text{VaR}_\alpha(\epsilon)}{\partial \epsilon} = -\mathbf{E}[Y | X + \epsilon Y = -\text{VaR}_\alpha(\epsilon)]. \quad (57)$$

The following proof is analogous to Gouriéroux, Laurent and Scaillet (2000). The mentioned authors have derived the expression for the derivative if existing, but have not proven the existence (of the derivative).

*Proof* (Gouriéroux, Laurent and Scaillet, 2000). If  $\partial \text{VaR}_\alpha(\epsilon)/\partial \epsilon$  exists, we have

$$\int \int_{-\infty}^{-\text{VaR}_\alpha(\epsilon) - \epsilon y} f(x, y) dx dy = \alpha, \quad (58)$$

and hence by differentiation with respect to  $\epsilon$

$$\int [-\partial \text{VaR}_\alpha(\epsilon)/\partial \epsilon - y] f(-\text{VaR}_\alpha(\epsilon) - \epsilon y, y) dy = 0. \quad (59)$$

This implies

$$\frac{\partial \text{VaR}_\alpha(\epsilon)}{\partial \epsilon} = -\frac{\int y f(-\text{VaR}_\alpha(\epsilon) - \epsilon y, y) dy}{\int f(-\text{VaR}_\alpha(\epsilon) - \epsilon y, y) dy} \quad (60)$$

and therefore (57).  $\square$

As already mentioned, the main problem in this reasoning is the missing proof of the differentiability of VaR. Also the strict positivity of the integral  $\int f(-\text{VaR}_\alpha(\epsilon) - \epsilon y, y) dy$  should be an important ingredient in an “exact” proof of the lemma. In the paper of Tasche (1999), there is given a sufficient condition, named (S), for VaR-differentiation. However, condition (S) is in the most cases not easy to prove (the normal distribution excluded) and differentiation may be possible even if (S) is not fulfilled.

**LEMMA A.2.** *Under certain strong assumptions on the density  $f$ , the Expected Shortfall  $ES_\alpha(\epsilon) := ES_\alpha(X + \epsilon Y)$  can be differentiable in  $\epsilon$  and*

$$\frac{\partial ES_\alpha(\epsilon)}{\partial \epsilon} = -\mathbf{E}[Y|X + \epsilon Y \leq -\text{VaR}_\alpha(\epsilon)]. \quad (61)$$

*Proof.* We have

$$ES_\alpha(\epsilon) = -\frac{1}{\alpha} \int \int_{-\infty}^{-\text{VaR}_\alpha(\epsilon) - \epsilon y} (x + \epsilon y) f(x, y) dx dy. \quad (62)$$

Differentiation with respect to  $\epsilon$  leads to

$$\begin{aligned} \frac{\partial ES_\alpha(\epsilon)}{\partial \epsilon} &= -\frac{1}{\alpha} \int \int_{-\infty}^{-\text{VaR}_\alpha(\epsilon) - \epsilon y} y f(x, y) dy \\ &\quad + \frac{1}{\alpha} \int \text{VaR}_\alpha(\epsilon) [-\partial \text{VaR}_\alpha(\epsilon) / \partial \epsilon - y] f(-\text{VaR}_\alpha(\epsilon) - \epsilon y, y) dy. \end{aligned} \quad (63)$$

Due to (60), the second summand is 0. □

In a more general context, Tasche (1999) also derives (61). Again, the most important parts of an existence proof would be the existence of the respective integrals and the proof of the correct application of all used differentiation rules.

## B Tables and figures

Share	$V_i(0)$	$\hat{\mu}_i$	$\hat{\sigma}_i$
Xetra DAX	2942.04	-0.54	0.45
Allianz	56.17	-1.46	0.78
BASF	38.16	-0.29	0.31
BMW	29.06	-0.50	0.32
Bayer	16.75	-0.80	0.66
Commerzbank	8.32	-0.92	0.79
DaimlerChrysler	28.90	-0.65	0.36
Deutsche Bank	44.86	-0.59	0.47
Lufthansa	8.79	-0.56	0.48
E.on	41.80	-0.31	0.29
Hypovereinsbank	10.20	-1.41	0.92

Table 1: Stock market parameters (1 year history). In the portfolio with 4 assets the Xetra DAX is treated like a single stock.

$\hat{a}_1$	$\hat{b}_1$	$\hat{\sigma}_1$	$\hat{\lambda}_1$
0.2648	0.0120	0.1236	-0.0647
$\hat{a}_2$	$\hat{b}_2$	$\hat{\sigma}_2$	$\hat{\lambda}_2$
1.7563	0.0145	0.1704	0.4968

Table 2: Estimates for the CIR-2-model (1 year history)

$$\begin{pmatrix} 1 & 0 & 0.7333 & 0.5860 \\ 0 & 1 & -0.4180 & -0.3799 \\ 0.7333 & -0.4180 & 1 & 0.9062 \\ 0.5860 & -0.3799 & 0.9062 & 1 \end{pmatrix}$$

Table 3: Adjusted covariance matrix  $\hat{\Sigma}$  (4 assets)

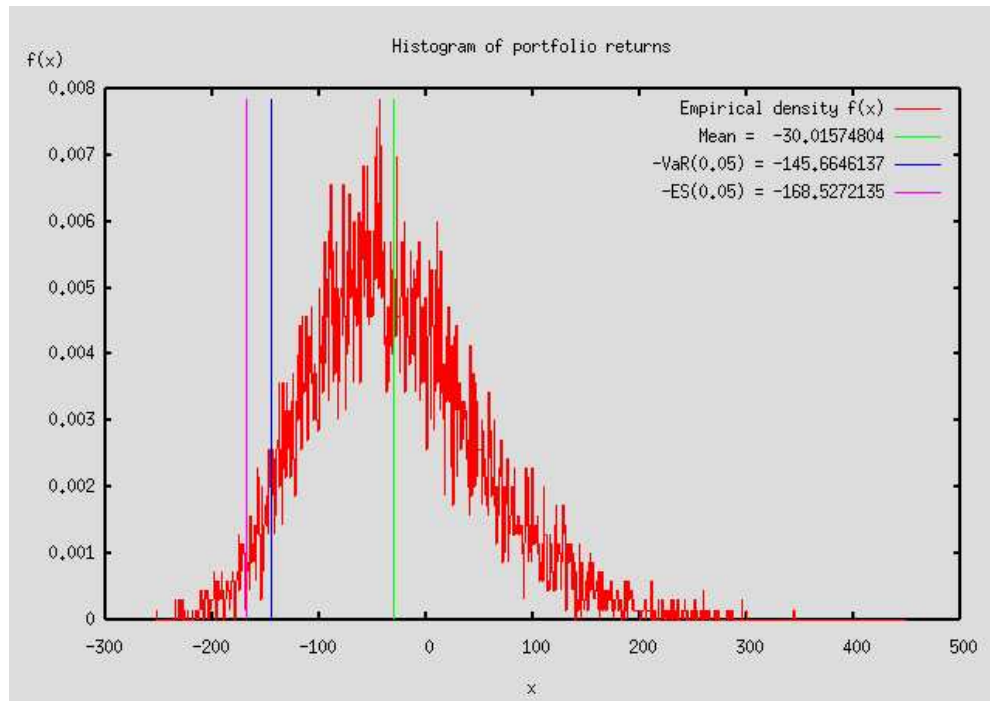


Figure 3: Histogram of returns; normed portfolio with 4 assets,  $10^4$  loops

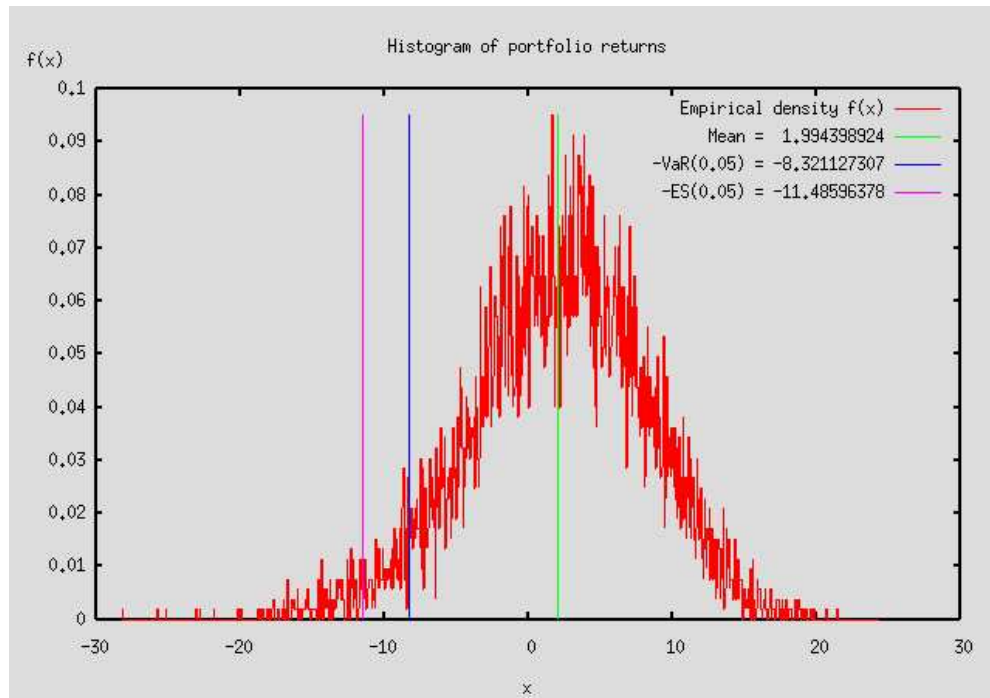


Figure 4: Histogram of returns; ES-optimized portfolio with 4 assets (local GS),  $10^4$  loops. Compared to the original portfolio in Figure 3, all values have tremendously been improved (please note the different scales).



Portfolio	Mean	VaR	ES	ES-RORC	ES-RORAC
normed Units Capital	-29.21	148.56	169.90	-0.1719	-0.0250
	(256.03, 382.88, 0.0850, 4.4508)				
	(250.0, 250.0, 250.0, 250.0)				
ES-opt. Units Capital	2.20	8.02	11.67	0.1888	0.0022
	(575.22, 595.47, 0.0284, -0.6059)				
	(561.67, 388.81, 83.54, -34.03)				
RORC-opt. Units Capital	8.73	18.61	28.39	0.3071	0.0085
	(506.74, 650.08, 0.0856, -3.0470)				
	(494.81, 424.47, 251.87, -171.15)				
RORAC-opt. Units Capital	14535.32	34795.68	56428.72	0.2576	0.2531
	(89585.98, -170200.78, 102.93, -4952.51)				
	(87476.85, -111132.65, 302838.23, -278182.42)				

Table 4: 4 assets;  $\alpha = 0.05$ ; no constraints; locally GS-optimized portfolios; GS started at normed portfolio

Portfolio	Mean	VaR	ES	ES-RORC	ES-RORAC
normed Units Capital	-29.21	148.56	169.90	-0.1719	-0.0250
	(256.03, 382.88, 0.0850, 4.4508)				
	(250.0, 250.0, 250.0, 250.0)				
ES-opt. Units Capital	2.75	1.11	2.19	1.2555	0.0027
	(1758.10, -1049.55, -0.0115, 0.0413)				
	(1716.71, -685.31, -33.73, 2.32)				
RORC-opt. Units Capital	3.07	0.84	1.96	1.5643	0.0031
	(1533.04, -730.19, -0.0036, -0.1729)				
	(1496.95, -476.78, -10.46, -9.71)				
RORAC-opt. Units Capital	29454.12	70484.98	114339.65	0.2576	0.2554
	(186074.79, -353548.05, 208.71, -10038.90)				
	(181693.00, -230849.31, 614040.41, -563885.10)				

Table 5: 4 assets;  $\alpha = 0.05$ ; no constraints; SI-GS-optimized portfolios

Portfolio	Mean	VaR	ES	ES-RORC	ES-RORAC
normed Units Capital	-29.21	148.56	169.90	-0.1719	-0.0250
	(256.03, 382.88, 0.0850, 4.4508)				
	(250.0, 250.0, 250.0, 250.0)				
ES-opt. Units Capital	1.81	4.19	6.26	0.2894	0.0018
	(1020.03, 0.22, 0.0013, 0.0001)				
	(996.01, 0.14, 3.84, 0.01)				
RORC-opt. Units Capital	1.88	4.29	6.37	0.2957	0.0019
	(1020.60, 4.22, 0.0000, 0.0119)				
	(996.57, 2.76, 0.01, 0.67)				
RORAC-opt. Units Capital	2.13	19.43	25.32	0.0843	0.0021
	(84.47, 1404.48, 0.0001, 0.0027)				
	(82.48, 917.06, 0.31, 0.15)				

Table 6: 4 assets;  $\alpha = 0.05$ ; constraint  $b = 0$ ; SI-GS-optimized portfolios

Portfolio	Mean	VaR	ES	ES-RORC	ES-RORAC
normed	-19.29	115.36	134.59	-0.1433	-0.0170
ES-opt.	0.32	6.93	8.87	0.0361	0.0003
RORC-opt.	7.13	8.28	13.79	0.5172	0.0070
RORAC-opt.	653.55	884.70	1369.21	0.4773	0.2759

Table 7: 20 assets;  $\alpha = 0.05$ ; no constraints; locally GS-optimized portfolios; GS started at normed portfolio

Portfolio	Mean	VaR	ES	ES-RORC	ES-RORAC
normed	-19.29	115.36	134.59	-0.1433	-0.0170
ES-opt.	2.52	2.23	3.56	0.7064	0.0025
RORC-opt.	4.40	2.92	4.82	0.9129	0.0044
RORAC-opt.	651.75	842.94	1357.35	0.4802	0.2765

Table 8: 20 assets;  $\alpha = 0.05$ ; no constraints; SI-GS-optimized portfolios

Portfolio	Mean	VaR	ES	ES-RORC	ES-RORAC
normed	-19.29	115.36	134.59	-0.1433	-0.0170
ES-opt.	-2.10	18.59	22.75	-0.0922	-0.0021
RORC-opt.	-2.22	23.76	27.84	-0.0799	-0.0021
RORAC-opt.	-2.1723	19.01	23.17	-0.0937	-0.0021

Table 9: 20 assets;  $\alpha = 0.05$ ; constraint  $b = 0$ ; SI-GS-optimized portfolios

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