

Quantum-inspired Evolutionary Algorithms for Calibration of the VG Option Pricing Model

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Abstract. Quantum effects are a natural phenomenon and just like evolution, or immune systems, can serve as an inspiration for the design of computing algorithms. This study illustrates how a quantum-inspired evolutionary algorithm can be constructed and examines the utility of the resulting algorithm on a problem in financial modelling known as model calibration. The results from the algorithm are shown to be robust and comparable to those of other algorithms.

1 Introduction

The objective of this study is to illustrate the potential for using a quantum rather than a traditional encoding representation in an evolutionary algorithm, and also to assess the utility of the resulting algorithm for the purposes of calibrating an option pricing model.

Quantum mechanics describes the behaviour of sub-atomic particles. Taking the example of an electron, it orbits the nucleus at different levels depending on its energy and its angular momentum. It can jump to a higher orbit by absorbing energy, or to a lower orbit by emitting energy. The orbits are discrete (or quantum levels) in that an electron cannot inhabit any in-between states or orbits. According to quantum mechanics, the location of electrons (or other subatomic particles) can best be represented by a quantum state vector. Broadly speaking, this vector describes the probabilities that an electron is occupying each of the possible discrete locations (orbits) which it could occupy. In the strange world of quantum mechanics, the particle is said to occupy *all* possible discrete locations at the same time but when observed, its quantum state collapses and it will be seen to occupy a distinct, discrete, state.

In recent years there has been a substantial interest in the theory and design of quantum computers, and the design of programs which could run on such computers. One interesting strand of research has been the use of natural computing (for example GP) to generate quantum circuits or programs (algorithms) for quantum computers [1]. There has also been associated work in a reverse direction which draws inspiration from concepts in quantum mechanics in order

to design novel natural computing algorithms. This is currently an area of active research interest. For example, quantum-inspired concepts have been applied to the domains of evolutionary algorithms [2–6], social computing [8], neuro-computing [9–11], and immuno-computing [12, 13]. A claimed benefit of these algorithms is that because they use a quantum representation, they can maintain a good balance between exploration and exploitation. It is also suggested that they offer computational efficiencies as use of a quantum representation can allow the use of smaller population sizes than typical evolutionary algorithms.

A particular potential benefit of applying a QEA methodology in the financial domain is

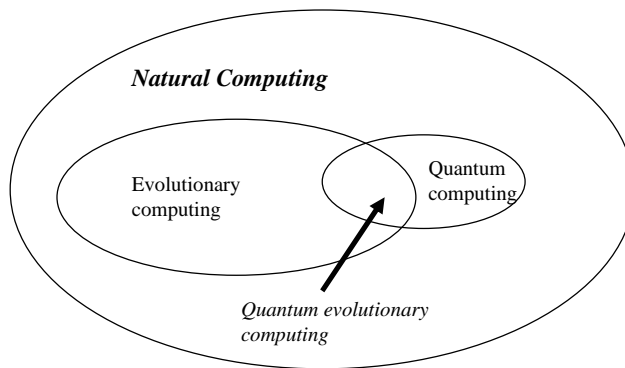


Fig. 1. Quantum-inspired evolutionary computing

Quantum-inspired algorithms offer interesting potential. As yet, due to their novelty, only a small number of recent papers have implemented a QEA, typically reporting good results [5, 6]. Consequently, we have a limited understanding of the performance of these algorithms and further testing is required in order to determine both their effectiveness and their efficiency. It is also noted that although a wide-variety of biologically-inspired algorithms have been applied for financial modelling [7], the QEA methodology has not yet been applied to the finance domain. This study addresses both of these gaps.

The rest of this paper is organised as follows. The next section provides a concise overview of QEA. We then outline the experimental methodology adopted. The remaining sections provide the results of these experiments followed by a number of conclusions.

2 The Quantum-inspired Genetic Algorithm

The best-known application of quantum-inspired concepts in evolutionary computing is the quantum-inspired genetic algorithm (QIGA) [2, 5, 6]. The (QIGA)

is based on the concepts of a qubit (quantum bit) and the superposition of states. In essence, in QIGAs the traditional representations used in evolutionary algorithms (binary, numeric and symbolic) are extended to include a quantum representation.

Under a quantum representation, the basic unit of information is no longer a bit which can assume two distinct states (0 or 1), but is a quantum system. Hence, a qubit (the smallest unit of information in a two-state quantum system) can assume either of the two ground states (0 or 1) or any superposition of the two ground states (the quantum superposition). A qubit can therefore be represented as

$$|q^i\rangle = \alpha|0\rangle + \beta|1\rangle \quad (1)$$

where $|0\rangle$ and $|1\rangle$ are the ground states 0 and 1, and α & β are complex numbers that specify the probability amplitudes of the two ground states. The act of observing (or measuring) a qubit projects the quantum system onto one of the ground states. $|\alpha|^2$ is the probability that the qubit will be in state 0 when it is observed, and $|\beta|^2$ is the probability that it will be in state 1. Hence, a qubit encodes the *probability* that a specific ground state will be seen when an observation takes place, rather than encoding the ground states themselves. In order to ensure this probabilistic interpretation remains valid, the values for α and β are constrained such that $|\alpha|^2 + |\beta|^2 = 1$.

More generally, a quantum system of m qubits can represent a total of 2^m states simultaneously. In the language of evolutionary computation a system of m qubits can be referred to as a *quantum chromosome* and can be written as a matrix

$$\begin{bmatrix} \alpha_1 & \alpha_2 & \dots & \alpha_m \\ \beta_1 & \beta_2 & \dots & \beta_m \end{bmatrix} \quad (2)$$

A key point when considering quantum systems is that they can compactly convey information on a large number of possible system states. In classical bit strings, a string of length n can represent 2^n possible states. However, a quantum space of n qubits has 2^n *dimensions*. This means that even a short qubit can convey information on many possible system states. For example, a 3 bit quantum system can encode 8 (2^3) distinct binary strings, and an 8 bit quantum system can encode 256 distinct strings.

Another important point in the context of applying quantum concepts to design evolutionary algorithms is that due to its probabilistic interpretation, a single qubit of length m can simultaneously represent **all** possible bit strings of length 2^m .

This implies that it is possible to modify standard evolutionary algorithms to work with a single quantum individual, rather than having to use a population of solution encodings. The qubit representation of the system states can also help maintain diversity during the search process of an evolutionary algorithm, due to its capability to represent multiple system states simultaneously.

2.1 Representing a Quantum System

There are many ways that a quantum system could be defined in order to encode a set of binary (solution) strings. For example, in the following 3 qubit quantum system, the quantum chromosome is defined using the three pairs of amplitudes below

$$\begin{bmatrix} \frac{1}{\sqrt{2}} & \frac{\sqrt{3}}{2} & \frac{1}{2} \\ \frac{1}{\sqrt{2}} & \frac{1}{2} & \frac{\sqrt{3}}{2} \end{bmatrix} \quad (3)$$

These probability amplitudes are the ratios that the particle (unit of information) is in a particular ground state rather than another. Hence, taking the first qubit, the occurrence of either state 0 or 1 is equally likely as both α_1 and β_1 have the same probability amplitude. Following on from the definition of the 3 qubit system, the state of the system is given by

$$\frac{\sqrt{3}}{4\sqrt{2}}|000\rangle + \frac{3}{4\sqrt{2}}|001\rangle + \frac{1}{4\sqrt{2}}|010\rangle + \frac{\sqrt{3}}{4\sqrt{2}}|011\rangle + \frac{\sqrt{3}}{4\sqrt{2}}|100\rangle + \frac{3}{4\sqrt{2}}|101\rangle + \frac{1}{4\sqrt{2}}|110\rangle + \frac{\sqrt{3}}{4\sqrt{2}}|111\rangle \quad (4)$$

To provide intuition on this point, consider the system state $|000\rangle$. The associated probability amplitude for this state is $\frac{\sqrt{3}}{4\sqrt{2}}$ and this is derived from the probability amplitudes of the 0 state for each of the three individual qubits ($\frac{1}{\sqrt{2}} * \frac{\sqrt{3}}{2} * \frac{1}{2} = 0.25$).

The associated probabilities of each of the individual states ($|000\rangle, |001\rangle, |010\rangle, |011\rangle, |100\rangle, |101\rangle, |110\rangle, |111\rangle$) are $\frac{3}{32}, \frac{9}{32}, \frac{1}{32}, \frac{3}{32}, \frac{3}{32}, \frac{9}{32}, \frac{1}{32}, \frac{3}{32}$ respectively. Taking the first of these states as an example, $\frac{\sqrt{3}}{4\sqrt{2}}^2 = \frac{3}{16}$.

2.2 The Algorithm

As already seen in previous chapters, a wide variety of algorithms can be derived using naturally-occurring phenomena as their inspiration. Similarly, there is no single QIGA, rather there are a family of possible algorithms which could be derived from the joint quantum-evolutionary metaphor. However, the following algorithm provides an example of a canonical QIGA

```

Set t=0
Initialise Q(t)
Create P(t) by undertaking an observation of Q(t)
Evaluate P(t) and select the best solution
Store the best solution in P(t) into B(t)
While (t < max t)
    t=t+1
    Create P*(t) by undertaking observations of Q(t-1)
    Evaluate P*(t)

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Update Q(t)
Store the best solutions in B(t-1) and P(t) into B(t)
Endwhile

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Initially, the population of quantum chromosomes is created

$Q(t) = q_1(t), q_2(t), \dots, q_n(t)$, where n is the population size, and each member of the population consists of an individual qubit of length m . The α and β values for each qubit are set to $\frac{1}{\sqrt{2}}$ in order to ensure that the states 0 and 1 are equally likely for each qubit.³ If there is domain knowledge that some states are likely to lead to better results, this can be used to seed the initial quantum chromosome(s).

Once a population of quantum chromosomes are created, these can be used to create a population of binary (or solution encoding) strings by performing an ‘observation’ on the quantum chromosomes. One way of performing the observation step is to draw a random number $rnd \in [0, 1]$. If $rnd > |\alpha_i(t)|^2$, the corresponding bit (j) in $p_i^j(t)$ is assigned state 1, otherwise it is assigned state 0.

Due to the stochastic nature of the observation step, the QIGA could be implemented using a single quantum chromosome, where this chromosome is observed multiple times in order to generate the population $P(t) = p_1(t), p_2(t), \dots, p_i(t)$. Alternatively, a small population of quantum chromosomes could be maintained, with each chromosome being observed a fixed number of times in order to generate $P(t)$.

In the while loop, an update step is performed on the quantum chromosome(s). This update step could be performed in a variety number of ways, for example by using pseudo-genetic operators, or by using a suitable quantum gate (see below). However the step is undertaken, its essence is that the quantum chromosome is adjusted in order to make the generation of the best solution found so far, more likely in the next iteration. As the optimal solution is approached by the QIGA system, the values of each element of the quantum chromosome tend towards either 0 or 1, corresponding to a high probability that the quantum chromosome will generate a specific solution vector (p_i) when observed.⁴

A number of distinctions between the QIGA above and the canonical GA (CGA) can be noted. In the CGA, the population of solutions persists from generation to generation, albeit in a changing form. In contrast, in QIGA, the population of solutions in $P(t)$ are discarded at the end of each loop. The described QIGA, unlike CGA, does not have explicit concepts of crossover or mutation. However, the adaptation of the quantum chromosomes in each iteration does embed implicit selection as the best solution is selected and is used to adapt the quantum chromosome(s). The crossover and mutation steps are also implicitly

³ The probability of either state 0 or 1 is $(\frac{1}{\sqrt{2}})^2 = 0.50$.

⁴ For example, a quantum chromosome (1 1 1) will generate an observed solution chromosome of (0 0 0) with probability 1, regardless of the choice of the parameter rnd in the observation step.

present, as the adaptation of the quantum chromosome in effect creates diversity, as it makes different states of the system more or less likely over time. Another distinction between the QIGA and the CGA is that the CGA operates directly on representations of the solution (the members of the current population of solutions), whereas in QIGA the update step is performed on the probability amplitudes of the ground states for each qubit making up the quantum chromosome(s).

2.3 Updating the Quantum Chromosome

Unlike the CGA, the QIGA does not typically employ distinct crossover and mutation operators. Instead a single diversity-generation step is used. There are two common approaches to this.

Quantum Mutation Quantum mutation is loosely inspired by the standard GA mutation operator. However, this is adapted so that the mutation step is guided by the best individual found to date, with the quantum chromosome being altered in order to make the generation of this solution more likely in future iterations of the algorithm [5, 6].

$$Q_{pointer}(t) = a * B_{bestsolution}(t) + (1 - a) * (1 - B_{bestsolution}(t)) \quad (5)$$

$$Q(t + 1) = Q_{pointer}(t) + b * randnorm(0, 1) \quad (6)$$

where $B_{bestsolution}(t)$ is the best solution found by iteration t . $Q_{pointer}(t)$ is a temporary quantum chromosome which is used to guide the generation of $Q(t+1)$ towards the form of $B_{bestsolution}$. The term $randnorm(0, 1)$ is a random number drawn from a (0,1) normal distribution. The parameters a and b control the balance between exploration and exploitation, with a governing the importance attached to $B_{bestsolution}(t)$ and b governing the degree of variance generation, centred on $Q_{pointer}(t)$. Values of $a \in [0.1, 0., 5]$ and $b \in [0.05, 0.15]$ are suggested by [5, 6].

Another simple method for generating diversity in the quantum chromosome [?] is as follows

$$\alpha(t + 1) = [2 - B_{bestsolution}(t)] * 0.5 \quad (7)$$

Under this mechanism, the bits in the current best individual will be more likely to be generated (their probability > 0.5) under and observation of the quantum chromosome in the next iteration of the algorithm.

Quantum Gates An alternative way to generate the quantum mutation is to use a transformation matrix (quantum rotation gate). The idea in using the quantum rotation gate is to steer the direction of the mutation towards the best performing individual in the population.

The i^{th} qubit value (α_i, β_i) is updated as follows

$$\begin{bmatrix} \alpha'_i \\ \beta'_i \end{bmatrix} = \begin{bmatrix} \cos(\theta_i) & -\sin(\theta_i) \\ \sin(\theta_i) & \cos(\theta_i) \end{bmatrix} \begin{bmatrix} \alpha_i \\ \beta_i \end{bmatrix} \quad (8)$$

where θ_i is the rotation angle of each qubit towards either 0 or 1. The definition of θ_i is problem specific and it controls the speed of convergence of the algorithm.

Readers are referred to [3] for more detail on this approach.

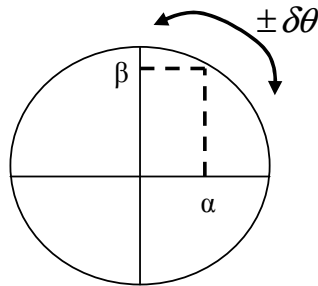


Fig. 2. Quantum mutation. The value $\pm\delta\theta$ is determined by reference to the value of the corresponding bit in the best solution found to date

3 Experimental Approach

An optimisation problem in financial modelling is considered to test the performance of the QIGA. The optimisation involves calibrating an option pricing model to observed market data. Calibration is a method of choosing model parameters so that the distance between a set of model option prices and market option prices is minimised, where distance is some metric such as the sum of squared errors or the sum of squared percentage errors. The parameters can be thought to resemble the markets view on current option prices and the underlying asset price. In calibration we do not explicitly take into account any historical data. All necessary information is contained in today's option prices which can be observed in the market. Practitioners frequently calibrate option pricing models so that the models provides a reasonable fit to current observed market option prices and they then use these models to price exotic derivatives

or for hedging purposes. In this paper we calibrate a popular extension of the Black-Scholes [16] option pricing model known as the Variance Gamma (*VG*) model [17–19] to FTSE 100 index option data.

A European call option on an asset S_t with maturity date T and strike price K is defined as a contingent claim with payoff at time T given by $\max[S_T - K, 0]$. The well known Black-Scholes (BS) formula for the price of a call on this asset is given by

$$C_{BS}(S_t, K, r, q, \tau; \sigma) = S_t e^{-q\tau} N(d_1) - K e^{-r\tau} N(d_2)$$

$$d_1 = \frac{-\ln m + (r - q + \frac{1}{2}\sigma^2)\tau}{\sigma\sqrt{\tau}} \quad d_2 = d_1 - \sigma\sqrt{\tau}$$

where $\tau = T - t$ is the time-to-maturity, t is the current time, $m = K/S$ is the moneyness of the option, r and q are the continuously compounded risk-free rate and dividend yield and $N(\cdot)$ is the cumulative normal distribution function. Suppose a market option price, denoted by $C_M(S_t, K)$, is observed. The Black-Scholes implied volatility for this option price is that value of volatility which equates the BS model price to the market option price as follows

$$\begin{aligned} \sigma_{BS}(S_t, K) &> 0 \\ C_{BS}(S_t, K, r, \tau; \sigma_{BS}(S_t, K)) &= C_M(S_t, K) \end{aligned}$$

If the assumptions underlying the BS option pricing model were correct, the BS implied volatilities for options on the same underlying asset would be constant for different strike prices and maturities. However in reality the BS implied volatilities are varying over strike price and maturity. Given that the options are written on a single underlying asset this result seems at first paradoxical, i.e. we have a number of different implied volatilities for a single asset which should only have one measure for its volatility. Yet if we relax some of the assumptions in the BS model, such as allowing for a more complex data generating process for the asset price than the log normal stochastic process (as assumed by BS), and take into account the resulting complications, this result begins to make sense and is simply highlighting the erroneous assumptions that underpin the BS model.

Many different option pricing models have been proposed as alternatives to the BS model. Examples include stochastic volatility models and jump diffusion models which allow for more complex asset price dynamics. We examine one such simple extension of the BS model known as the Variance Gamma (*VG*) option pricing model. The idea is to model stock price movements occurring on business time rather than on calendar time using a time transformation of a Brownian motion. The resulting model is a three parameter model where roughly speaking we can interpret the parameters as controlling volatility, skewness and kurtosis, denoted respectively as σ, θ and ν , of the underlying asset returns distribution.

Closed form option pricing formulae exist under the *VG* model [19].

$$C_{VG}(S_t, K, r, \tau; \{\sigma, \nu, \theta\}) = S_t e^{-q\tau} \Psi \left(d \sqrt{\frac{1-c_1}{\nu}}, (\alpha + s) \sqrt{\frac{\nu}{1-c_1}}, \frac{\tau}{\nu} \right) - K e^{-r\tau} \Psi \left(d \sqrt{\frac{1-c_2}{\nu}}, \alpha s \sqrt{\frac{\nu}{1-c_2}}, \frac{\tau}{\nu} \right)$$

where

$$d = \frac{1}{s} \left[\ln \left(\frac{S_t}{K} \right) + (r - q) \tau + \frac{\tau}{\nu} \ln \left(\frac{1-c_1}{1-c_2} \right) \right]$$

$$\alpha = \zeta s, \quad \zeta = -\frac{\theta}{\sigma^2}, \quad s = \frac{\sigma}{\sqrt{1 + \left(\frac{\theta}{\sigma}\right)^2 \frac{\nu}{2}}}$$

$$c_1 = \frac{\nu(\alpha + s)^2}{2}, \quad c_2 = \frac{\nu\alpha^2}{2}$$

and where Ψ is defined in terms of the modified Bessel function of the second kind⁵.

4 Results

Market makers in the options markets quote BS implied volatilities rather than option prices even though they realise BS is a flawed model. The first row in Table 1 depicts end-of-day settlement Black-Scholes implied volatilities for FTSE 100 European options on the 17 March 2006 for different strike prices and a time-to-maturity of 35 days. As can be seen the BS implied volatilities are not constant across the strike price. The second and third row in Table 1 converts the BS implied volatilities into market call and put prices by substituting the BS implied volatilities into the Black-Scholes formula. The following input parameters were used to calculate the option prices, the index price is the FTSE 100 index itself $S_t = 5999.4$, the interest rate is the one month Libor rate converted into a continuously compounded rate $r = 0.0452$ and the dividend yield is a continuously compounded dividend yield downloaded from Datastream and is $q = 0.0306$. These prices are then taken to be the observed market option prices. Out-of-the money (OTM) option prices are considered most suitable for calibration purposes because of their liquidity and informational content. Hence OTM call prices were used for $K < S$ and OTM put prices were used for $K > S$ in the calibration. The calibration problem now amounts to choosing an optimum parameter vector $\Theta = \{\sigma, \nu, \theta\}$ such that an objective function $G(\Theta)$ is minimised. In this paper the objective function is chosen to be the absolute average percentage error (APE)

$$G(\Theta) = \frac{1}{N} \sum_{i=1}^N \left| \frac{C_i - C_i(\Theta)}{C_i} \right|$$

⁵ See [19] for exact form of this function.

where C_i is the observed market price on the i -th option (could be a call or a put) and $C_i(\Theta)$ is the VG model price of the i -th option with parameter vector Θ . One of the difficulties in model calibration is that the available market information may be insufficient to completely identify the parameters of a model [20]. If the model is sufficiently rich relative to the number of market prices available a number of possible parameter vector combinations will be compatible with market prices and the objective function $G(\Theta)$ may not be convex function of Θ . A plot of the objective function versus the two parameters controlling skewness and kurtosis of the asset returns distribution, θ and ν , whilst keeping σ fixed at $\sigma = 0.1116$ is shown in figure 3(a).

Strike price	5695.2	5845.1	5995.0	6144.9	6294.7
IV (%)	13.76	12.41	11.13	10.44	10.94
Call(\$)	323.67	193.63	88.67	28.03	7.99
Put (\$)	12.44	31.63	75.89	164.48	293.67

Table 1. Market BS implied volatilities and option prices for FTSE 100 index options on the 17 March 2006. The strike prices are given in the table and the other observable inputs are $S = 5999.4$, $\tau = \frac{35}{365}$, $r = 0.0452$ and $q = 0.0306$.

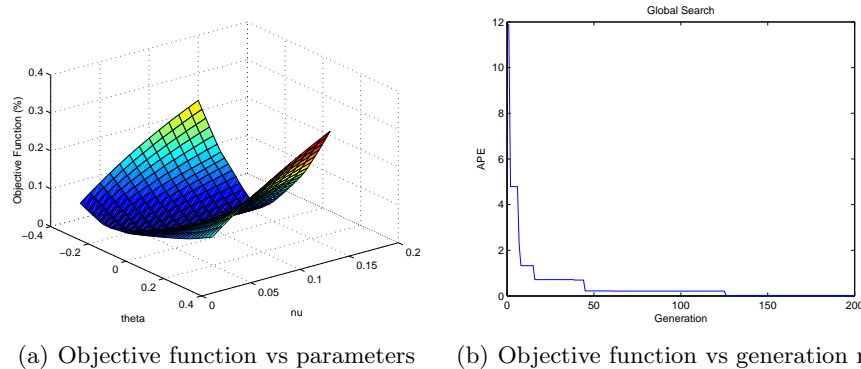


Fig. 3. Objective function versus model parameters ν and θ and objective function versus generation number.

It displays a flat profile near the minimum where many parameter combinations will yield equivalent fits. The error surface is not a straightforward error surface and a local optimiser might not converge to the true optimum. There are regions where the error surface is very flat for changes in the parameter val-

ues and there are regions where the optimiser might get not converge to global optimum.

4.1 Comparison of Results

To provide a benchmark for the results obtained by the QIGA a deterministic matlab optimiser called `fminsearch` was run 20 times with different initial parameter vectors. The matlab optimiser converged to different values for Θ for different initialisations of the parameter vector so the one with the optimal value for the objective function G was chosen. The results are reported in the Tables 2 and 3. As can be seen when averaged over only 20 runs the QIGA parameter vector Θ is very close to the optimal parameter vector from matlab. Figure 3(b) depicts the evolution of the global objective function G (also known as APE) as a function of the generation number. Figures 4(a) and 4(b) depict the evolution of the parameters ν and θ as a function of the generation number. The results reported in Table 4 are where the exploitation and exploration parameters are changed to those values reported in the table and everything else remains fixed. It can be seen that the performance of the QIGA is not as good as with the previous values and perhaps the exploitation parameter is too small and the exploration parameter is too large for this problem. Further sensitivity analysis would need to be conducted to find optimal values for these parameters.

Parameter setting	
number of runs	20
generation number	200
population size	10
exploitation	0.3
exploration	0.5

Table 2. Parameter settings that were used in the QIGA.

Parameter	QIGA	Matlab	Market Price	Mean Model Price	Mean APE
mean σ	0.1138	0.1143	12.44	12.44	0.08056
mean ν	0.0667	0.0638	31.64	31.55	
mean θ	-0.1396	-0.1429	75.90	75.56	
			28.02	28.29	
			7.99	8.03	

Table 3. Results of QIGA where the average parameter values after 20 runs are compared with the average parameters from 20 runs of a matlab optimiser. The resulting mean model prices from the 20 runs are compared with the market prices and the mean APE is reported.

Parameter	QIGA	Matlab	Market Price	Mean Model Price	Mean APE
mean σ	0.1137	0.1143	12.44	12.38	0.1443
mean ν	0.0540	0.0638	31.64	31.50	
mean θ	-0.1320	-0.1429	75.90	75.60	
			28.02	28.34	
			7.99	8.04	

Table 4. Results of QIGA where the average parameter values after 20 runs are compared with the average parameters from 20 runs of a matlab optimiser. The resulting mean model prices from the 20 runs are compared with the market prices and the mean APE is reported. The exploitation and exploration parameters have now been changed to 0.2ϵ and 0.6ϵ .

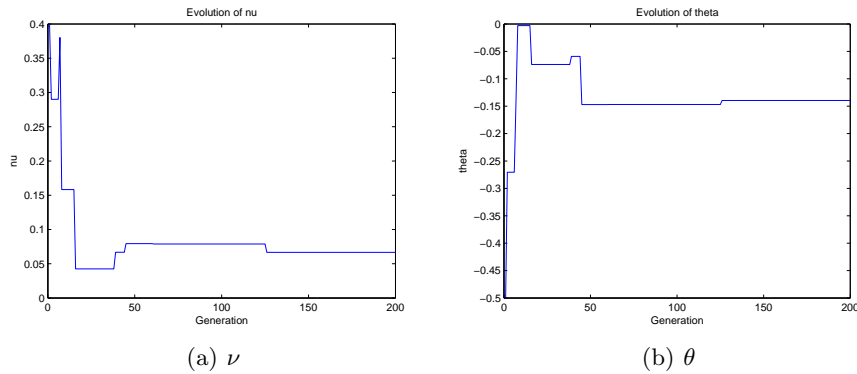


Fig. 4. Evolution of parameters ν and θ as a function of the generation number.

5 Conclusions & Future Work

Several extensions of the methodology in this study are indicated for future work. The first extension would be to extend the QIGA to a higher dimensional setting so that the computational benefits of QIGA really begin to take affect. Other extensions would include an analysis of the Q-gate operator and how it is updated through $\delta\theta$, and an analysis of the parameters used in the algorithm. Financial applications include the calibration and estimation of more complex higher dimensional models to market data in an evolutionary setting so that information on model uncertainty is calculated as an integral part of the algorithm. The use of QIGA in high dimensional financial modelling settings may be crucial due to the potential reduction in computational time.

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