Monte Carlo methods for security pricing

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Abstract

The Monte Carlo approach has proved to be a valuable and flexible computational tool in modern finance. This paper discusses some of the recent applications of the Monte Carlo method to security pricing problems, with emphasis on improvements in efficiency. We first review some variance reduction methods that have proved useful in finance. Then we describe the use of deterministic low-discrepancy sequences, also known as quasi-Monte Carlo methods, for the valuation of complex derivative securities. We summarize some recent applications of the Monte Carlo method to the estimation of partial derivatives or risk sensitivities and to the valuation of American options. We conclude by mentioning other applications.

Keywords: Monte Carlo simulation; Quasi-Monte Carlo; Option pricing; Variance reduction; & Derivative estimation

1. Introduction

In recent years the complexity of numerical computation in financial theory and practice has increased enormously, putting more demands on computational speed and efficiency. Numerical methods are used for a variety of purposes of finance. These include the valuation of securities, the estimation of their sensitivities, risk analysis, and stress testing of portfolios. The Monte Carlo method is a useful tool for many of these calculations, evidenced in part by the

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voluminous literature of successful applications. For a brief sampling, the reader is referred to the stochastic volatility applications in Duan (1995), Hull and White (1987), Johnson and Shanno (1987), and Scott (1987); the valuation of mortgage-backed securities in Schwartz and Torous (1989); the valuation of path-dependent options in Kemna and Vorst (1990); the portfolio optimization in Worzel et al. (1994); and the valuation of interest-rate derivative claims in Carverhill and Pang (1995). In this paper we focus on recent methodological developments. We review the Monte Carlo approach and describe some recent applications in the finance area.

In modern finance, the prices of the basic securities and the underlying state variables are often modelled as continuous-time stochastic processes. A derivative security, such as a call option, is a security whose payoff depends on one or more of the basic securities. Using the assumption of no arbitrage, financial economists have shown that the price of a generic derivative security can be expressed as the expected value of its discounted payouts. This expectation is taken with respect to a transformation of the original probability measure known as the equivalent martingale measure or the risk-neutral measure. The book by Duffie (1996) provides an excellent account of this material.

The Monte Carlo method lends itself naturally to the evaluation of security prices represented as expectations. Generically, the approach consists of the following steps:

- Simulate sample paths of the underlying state variables (e.g., underlying asset prices and interest rates) over the relevant time horizon. Stimulate these according to the risk-neutral measure.
- Evaluate the discounted cash flows of a security on each sample path, as determined by the structure of the security in question.
- Average the discounted cash flows over sample paths.

In effect, this method computes a multi-dimensional integral—the expected value of the discounted payouts over the space of sample paths. The increase in the complexity of derivative securities in recent years has led to a need to evaluate high-dimensional integrals.

Monte Carlo becomes increasingly attractive compared to other methods of numerical integration as the dimension of the problem increases. Consider the integral of the function \( f(x) \) over the \( d \)-dimensional unit hypercube. The simple (or crude) Monte Carlo estimate of the integral is equal to the average value of the function \( f \) over \( n \) points selected at random\(^2\) from the unit hypercube. From the strong law of large numbers this estimate converges to the true value of the

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\(^1\) Wiggins (1987) also studies pricing under stochastic volatility but does not use Monte Carlo simulation.

\(^2\) In standard Monte Carlo application the \( n \) points are usually not truly random but are generated by a deterministic algorithm and are described as pseudorandom numbers.
integrands as $n$ tends to infinity. In addition, the central limit theorem assures us that
the standard error$^3$ of the estimate tends to zero as $1/\sqrt{n}$. Thus, the error
convergence rate is independent of the dimension of the problem and this is the
dominant advantage of the method over classical numerical integration
approaches. The only restriction on the function $f$ is that it should be square
integrable, and this is a relatively mild restriction.

Furthermore, the Monte Carlo method is flexible and easy to implement and
modify. In addition, the increased availability of powerful computers has
enhanced the attractiveness of the method. There are some disadvantages of
the method but in recent years progress has been made in overcoming them.
One drawback is that for very complex problems a large number of replications
may be required to obtain precise results. Different variance reduction
techniques have been developed to increase precision. Two of the classical
variance reduction techniques are the control variate approach and the
antithetic variate method. More recently, moment matching, importance
sampling, and conditional Monte Carlo methods have been introduced in finance
applications.

Another technique for speeding up the valuation of multi-dimensional integrals
uses deterministic sequences rather than random sequences. These
deterministic sequences are chosen to be more evenly dispersed throughout the
region of integration than random sequences. If we use these sequences to
estimate multi-dimensional integrals we can often improve the convergence.
Deterministic sequences with this property are known as low-discrepancy se-
quences or quasi-random sequences. Using this approach one can in theory
derive deterministic error bounds, though the practical use of the bounds is
problematic. In contrast, standard Monte Carlo yields simple, useful probabilis-
tic error bounds. Although low-discrepancy sequences are well known in com-
putational physics they have only recently been applied in finance problems.
There are different procedures for generating such low-discrepancy sequences
and these procedures are generally based on number theoretic methods. We
describe some of the recent developments in this area. We also discuss applica-
tions of this approach to problems in finance and conduct some rough compar-
isons between standard Monte Carlo methods and two different quasi-random
approaches.

Until recently, the valuation of American style options was widely considered
outside the scope of Monte Carlo. However, Tilley (1993), Broadie and Glasser-
man (1997), and Barraquand and Martineau (1995) have proposed approaches
to this problem, and there has been other related work as well. We provide
a brief survey of the recent research progress in this area.

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$^3$ We can readily estimate the variance of the Monte Carlo estimate by using the same set of
$n$ random numbers to estimate the expected value of $f^2$. 
The layout of the paper is as follows. Variance reduction techniques are described in the next section. The ideas behind the use of low-discrepancy sequences and brief numerical comparisons with standard Monte Carlo methods are given in Section 3. Price sensitivity estimation using simulation is discussed in Section 4. Various approaches to pricing American options using simulation are briefly described in Section 5. Other issues are touched on briefly in Section 6.

2. Variance reduction techniques

In this section, we first discuss the role of variance reduction in meeting the broader objective of improving the computational efficiency of Monte Carlo simulations. We then discuss specific variance reduction techniques and illustrate their application to pricing problems.

2.1. Variance reduction and efficiency improvement

The reduction of variance seems so obviously desirable that the precise argument for its benefit is sometimes overlooked. We briefly review the underlying justification for variance reduction and examine it from the perspective of improving computational efficiency.

Suppose we want to compute a parameter \( \theta \) — for example, the price of a derivative security. Suppose we can generate by Monte Carlo an i.i.d. sequence \( \{ \hat{\theta}_i, i = 1, 2, \ldots \} \), where each \( \hat{\theta}_i \) has expectation \( \theta \) and variance \( \sigma^2 \). A natural estimator of \( \theta \) based on \( n \) replications is then the sample mean

\[
\frac{1}{n} \sum_{i=1}^{n} \hat{\theta}_i.
\]

By the central limit theorem, for large \( n \) this sample mean is approximately normally distributed with mean \( \theta \) and variance \( \sigma^2/n \). Probabilistic error bounds in the form of confidence intervals follow readily from the normal approximation, and indicate that the error in the estimator is proportional to \( \sigma/\sqrt{n} \). Thus, decreasing the variance \( \sigma^2 \) by a factor of 10, say, while leaving everything else unchanged, does as much for error reduction as increasing the number of samples by a factor of 100.

Suppose, now, that we have a choice between two types of Monte Carlo estimates which we denote by \( \{ \hat{\theta}_i^{(1)}, i = 1, 2, \ldots \} \) and \( \{ \hat{\theta}_i^{(2)}, i = 1, 2, \ldots \} \). Suppose that both are unbiased, so that \( \text{E}[\hat{\theta}_i^{(1)}] = \text{E}[\hat{\theta}_i^{(2)}] = \theta \), but \( \sigma_1 < \sigma_2 \), where \( \sigma_j^2 = \text{Var}[\hat{\theta}_j^{(j)}], j = 1, 2 \). From our previous observations it follows that a sample mean of \( n \) replications of \( \hat{\theta}_i^{(1)} \) gives a more precise estimate of \( \theta \) than does
a sample mean of $n$ replications of $\hat{\theta}^{(2)}$. But this analysis oversimplifies the
comparison because it fails to capture possible differences in the computational
effort required by the two estimators. Generating $n$ replications of $\hat{\theta}^{(1)}$ may be
more time-consuming than generating $n$ replications of $\hat{\theta}^{(2)}$; smaller variance is
not sufficient grounds for preferring one estimator over another.

To compare estimators with different computational requirements as well as
different variances, we argue as follows. Suppose the work required to generate
one replication of $\hat{\theta}^{(j)}$ is a constant $b_j, j = 1, 2$. (In some problems, the work per
replication is stochastic; assuming it is constant simplifies the discussion.) With
computing time $t$, the number of replications of $\hat{\theta}^{(j)}$ that can be generated is
$\lfloor t/b_j \rfloor$; for simplicity, we drop the $\lfloor \cdot \rfloor$ and treat the ratios $t/b_j$ as though they
were integers. The two estimators available with computing time $t$ are, therefore,

$$\frac{b_1}{t} \sum_{i=1}^{\lfloor t/b_1 \rfloor} \hat{\theta}_i^{(1)} \quad \text{and} \quad \frac{b_2}{t} \sum_{i=1}^{\lfloor t/b_2 \rfloor} \hat{\theta}_i^{(2)}.$$

For large $t$, these are approximately normally distributed with mean $\theta$ and with
standard deviations

$$\sigma_1 \sqrt{\frac{b_1}{t}} \quad \text{and} \quad \sigma_2 \sqrt{\frac{b_2}{t}}.$$

Thus, for large $t$, the first estimator should be preferred over the second if

$$\sigma_1^2 b_1 < \sigma_2^2 b_2. \quad (1)$$

Eq. (1) provides a sound basis for trading-off estimator variance and computa-
tional requirements. In light of the discussion leading to (1), it is reasonable
to take the product of variance and work per run as a measure of efficiency. Using efficiency as a basis for comparison, the lower-variance estimator should be
preferred only if the variance ratio $\sigma_1^2 / \sigma_2^2$ is smaller than the work ratio $b_2/b_1$.
By the same argument, a higher-variance estimator may actually be preferable if
it takes much less time to generate.

In its simplest form, the principle expressed in (1) dates at least to Hammersley
and Handscomb (1964, p. 22). More recently, the idea has been substantially
extended by Glynn and Whitt (1992). They allow the work per run to be random
(in which case each $b_j$ is the expected work per run) and also consider efficiency
in the presence of bias.

2.2. Antithetic variates

Equipped with a basis for evaluating potential efficiency improvements, we
can now consider specific variance reduction techniques. One of the simplest
and most widely used techniques in financial pricing problems is the method of antithetic variates. We introduce it with a simple example, then generalize.

Consider the problem of computing the Black-Scholes price of a European call option on a no-dividend stock. Of course, there is no need to evaluate this price by simulation, but the example serves as a useful introduction. In the Black-Scholes model, the stock price follows a log-normal diffusion. Independent replications of the terminal stock price under the risk-neutral measure can be generated from the formula

\[ S_T^{(i)} = S_0 e^{(-r + \sigma^2/2)T + \sigma \sqrt{T} \epsilon_i}, \quad i = 1, \ldots, n, \]  

where \( S_0 \) is the current stock price, \( r \) is the riskless interest rate, \( \sigma \) is the stock’s volatility, \( T \) is the option’s maturity, and the \( \{ \epsilon_i \} \) are independent samples from the standard normal distribution. See, e.g., Hull (1997) for background on this model, and see Devroye (1986) for methods of sampling from the normal distribution. Based on \( n \) replications, an unbiased estimator of the price of an option with strike \( K \) is given by

\[ \hat{C} = \frac{1}{n} \sum_{i=1}^{n} C_i \equiv \frac{1}{n} \sum_{i=1}^{n} e^{-rT} \max \{0, S_T^{(i)} - K\}. \]  

In this context, the method of antithetic variates is based on the observation that if \( Z_t \) has a standard normal distribution, then so does \( -Z_t \). The price \( S_T^{(i)} \) obtained from (2) with \( Z_t \) replaced by \( -Z_t \) is thus a valid sample from the terminal stock price distribution. Similarly, each

\[ \tilde{C}_i = e^{-rT} \max \{0, S_T^{(i)} - K\} \]

is an unbiased estimator of the option price, as is, therefore,

\[ \hat{C}_{AV} = \frac{1}{n} \sum_{i=1}^{n} C_i + \tilde{C}_i. \]

A heuristic argument for preferring \( \hat{C}_{AV} \) notes that the random inputs obtained from the collection of antithetic pairs \( \{(Z_i, -Z_i)\} \) are more regularly distributed than a collection of \( 2n \) independent samples. In particular, the sample mean over the antithetic pairs always equals the population mean of 0, whereas the mean over finitely many independent samples is almost surely different from 0. If the inputs are made more regular, it may be hoped that the

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4 This method was introduced to option pricing in Boyle (1977), where its use was illustrated in the pricing of a European call on a dividend-paying stock.
outputs are more regular as well. Indeed, a large value of $S_T^{(i)}$ resulting from a large $Z_i$ will be paired with a small value of $S_T^{(i)}$ obtained from $-Z_i$.

A more precise argument compares efficiencies. Because $C_i$ and $\tilde{C}_i$ have the same variance,

$$\text{Var} \left[ \frac{C_i + \tilde{C}_i}{2} \right] = \frac{1}{2} (\text{Var}[C_i] + \text{Cov}[C_i, \tilde{C}_i]).$$

(4)

Thus, we have $\text{Var}[\hat{C}_{AV}] \leq \text{Var}[\hat{C}]$ if $\text{Cov}[C_i, \tilde{C}_i] \leq \text{Var}[C_i]$. However, $\hat{C}_{AV}$ uses twice as many replications as $\hat{C}$, so we must account for differences in computational requirements. If generating the $Z_i$ takes a negligible fraction of the work per replication (which would typically be the case in the pricing of a more elaborate option), then the work to generate $\hat{C}_{AV}$ is roughly double the work to generate $\hat{C}$. Thus, for antithetics to increase efficiency, we require

$$2\text{Var}[\hat{C}_{AV}] \leq \text{Var}[\hat{C}],$$

which, in light of (4), simplifies to the requirement that $\text{Cov}[C_i, \tilde{C}_i] \leq 0$.

That this condition is met is easily demonstrated. Define $\phi$ so that $C_i = \phi(Z_i)$; $\phi$ is the composition of the mappings from $Z_i$ to the stock price and from the stock price to the discounted option payoff. As the composition of two increasing functions, $\phi$ is monotone, so by a standard inequality (e.g., Section 2.2 of Barlow and Proschan, 1975)

$$\mathbb{E}[\phi(Z_i)\phi(-Z_i)] \leq \mathbb{E}[\phi(Z_i)]\mathbb{E}[\phi(-Z_i)],$$

(5)

i.e., $\text{Cov}[C_i, \tilde{C}_i] = \mathbb{E}[\phi(Z_i)\phi(-Z_i)] - \mathbb{E}[\phi(Z_i)]\mathbb{E}[\phi(-Z_i)] \leq 0$, and we may conclude that antithetics help.

This argument can be adapted to show that the method of antithetic variates increases efficiency in pricing a European put and other options that depend monotonically on inputs (e.g., Asian options). The notable departure from monotonicity in some barrier options (e.g., a down-and-in call) suggests that the use of antithetics in pricing these options may sometimes be less effective.

In computing confidence intervals with antithetic variates, it is essential that the standard error be estimated using the sample standard deviation of the $n$ averaged pairs $(C_i + \tilde{C}_i)/2$ and not the $2n$ individual observations $C_1, \tilde{C}_1, \ldots, C_n, \tilde{C}_n$. The averaged pairs are independent but the individual observations are not. This is a case (we will see others shortly) in which the use of a variance reduction technique affects the estimation of the standard error and, in particular, requires some 'batching' of observations to deal with dependence.

It is worth noting that the method of antithetic variates is by no means restricted to simulations whose only stochastic inputs are standard normal
variates. The most primitive stochastic input in most simulations is a sequence \( \{ U_n \} \) of independent variates uniformly distributed on the unit interval. In this case, \( 1 - U_n \) has the same distribution as \( U_n \), and the pair \( (U_n, 1 - U_n) \) are called antithetic because they exhibit negative dependence. If the simulation output depends monotonically on the input random numbers, then the output obtained from \( \{ 1 - U_1, 1 - U_2, \ldots \} \) will be negatively correlated with that obtained from \( \{ U_1, U_2, \ldots \} \), resulting in increased efficiency compared with independent replications.

For further general background on antithetic variates and other methods based on correlation induction, see Bratley et al. (1987), Hammersley and Handscomb (1964), Glynn and Iglehart (1988), and references therein. For some examples of application in finance, see Boyle (1977), Clewlow and Carverhill (1994), and Hull and White (1987).

### 2.3. Control variates

The method of control variates is among the most widely applicable, easiest to use, and effective of the variance reduction techniques.\(^5\) Simply put, the principle underlying this technique is 'use what you know'.

The most straightforward implementation of control variates replaces the evaluation of an unknown expectation with the evaluation of the difference between the unknown quantity and another expectation whose value is known. A specific illustration can be found in the analysis of Boyle and Emanuel (1985) and Kemna and Vorst (1990) of Asian options. Let \( P_A \) be the price of an option whose payoff depends on the arithmetic average of the underlying asset. Let \( P_G \) be the price of an option equivalent in every respect except that a geometric average replaces the arithmetic average. Most options based on averages use arithmetic averaging, so \( P_A \) is of much greater practical value; but whereas \( P_A \) is analytically intractable, \( P_G \) can often be evaluated in closed form. Can knowledge of \( P_G \) be leveraged to compute \( P_A \)?

It can, through the control variate method. Write \( P_A = E[\hat{P}_A] \) and \( P_G = E[\hat{P}_G] \), where \( \hat{P}_A \) and \( \hat{P}_G \) are the discounted option payoffs for a single simulated path of the underlying asset. Then

\[
P_A = P_G + E[\hat{P}_A - \hat{P}_G];
\]

in other words, \( P_A \) can be expressed as the known price \( P_G \) plus the expected difference between \( \hat{P}_A \) and \( \hat{P}_G \). An unbiased estimator of \( P_A \) is thus provided by

\[
\hat{P}_A^{cv} = \hat{P}_A + (P_G - \hat{P}_G).
\]  

\(^5\) The earliest application of this technique to option pricing is Boyle (1977).
This representation suggests a slightly different interpretation; \( \hat{P}_A^a \) adjusts the straightforward estimator \( \hat{P}_A \) according to the difference between the known value \( P_G \) and the observed value \( \hat{P}_G \). The known error \( (P_G - \hat{P}_G) \) is used as a control in the estimation of \( P_A \).

If most of the computational effort goes to generating paths of the underlying asset, then the additional work required to evaluate \( \hat{P}_G \) along with \( \hat{P}_A \) is minor. It therefore seems reasonable to compare variances alone. Since

\[
\text{Var}[\hat{P}_A^a] = \text{Var}[\hat{P}_A] + \text{Var}[\hat{P}_G] - 2\text{Cov}[\hat{P}_A, \hat{P}_G],
\]

this method is effective if the covariance between \( \hat{P}_A \) and \( \hat{P}_G \) is large. The numerical results of Kemna and Vorst indicate that this is indeed the case. Fu, Madan, and Wang (1995) have investigated the use of other control variates for Asian options, based on Laplace transform values. These appear to be less strongly correlated with the option price.

A closer examination of (6) reveals that this estimator does not make optimal use of the relation between the two option prices. Consider the family of unbiased estimators

\[
\hat{P}_A^\beta = \hat{P}_A + \beta(P_G - \hat{P}_G),
\]

parameterized by the scalar \( \beta \). We have

\[
\text{Var}[\hat{P}_A^\beta] = \text{Var}[\hat{P}_A] + \beta^2 \text{Var}[\hat{P}_G] - 2\beta \text{cov}[\hat{P}_A, \hat{P}_G].
\]

The variance-minimizing \( \beta \) is, therefore,

\[
\beta^* = \frac{\text{Cov}[\hat{P}_A, \hat{P}_G]}{\text{Var}[\hat{P}_G]}.
\]

Depending on the application, \( \beta^* \) may or may not be close to 1, the implicit value in (6). In using an estimator of the form (6), we forgo an opportunity for greater variance reduction. Indeed, whereas (6) may increase or decrease variance, an estimator based on \( \beta^* \) is guaranteed not to increase variance, and will result in a strict decrease in variance so long as \( \hat{P}_A \) and \( \hat{P}_G \) are not uncorrelated.

In practice, of course, we rarely know \( \beta^* \) because we rarely know \( \text{Cov}[\hat{P}_A, \hat{P}_G] \). However, given \( n \) independent replications \( \{(P_{Ai}, \hat{P}_{Gi}) : i = 1, \ldots, n \} \) of the pairs \( (\hat{P}_A, \hat{P}_G) \) we can estimate \( \beta^* \) via regression. At this point

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*To go from (6) to Boyle's (1977) example, let \( P_G \) be the price of a European call option on a no-dividend stock and let \( P_A \) be the corresponding option price in the presence of dividends.*
we face a choice. Using all \( n \) replications to compute an estimate \( \hat{\beta} \) of \( \beta^* \) introduces a bias in the estimator

\[
\frac{1}{n} \sum_{i=1}^{n} P_{AI} + \hat{\beta}\left(P_G - \frac{1}{n} \sum_{i=1}^{n} P_{Gi}\right),
\]

and its estimated standard error because of the dependence between \( \hat{\beta} \) and the \( P_{Gi} \). Reserving \( n_1 \) replications for the estimation of \( \beta^* \) and the remaining \( n - n_1 \) replications for the sample mean of the \( P_{Gi} \) (typically with \( n_1 \ll n \)) eliminates the bias but may deteriorate the estimate of \( \beta^* \). Neither issue significantly limits the applicability of the method, because the possible bias vanishes as \( n \) increases and because the estimator of \( \beta^* \) need not be very precise to achieve a reduction in variance.

The advantage of working with (7) over (6) becomes even more pronounced when further controls are introduced. For example, when the asset price is simulated under risk-neutral probabilities, the present value \( e^{-rT}E[S_T] \) of the terminal price must equal the current price \( S_0 \). We can, therefore, form the estimator

\[
\hat{P}_A + \beta_1(P_G - \hat{P}_G) + \beta_2(S_0 - e^{-rT}S_T).
\]

The variance-minimizing coefficients \( (\beta_1^*, \beta_2^*) \) are easily found by multiple regression. This optimization step seems particularly crucial in this case; for whereas one might guess that \( \beta_1^* \) is close to 1, it seems unlikely that \( \beta_2^* \) would be. Optimizing over the \( \beta_2 \) allows us to exploit controls that are negatively correlated with the option payoff.


2.4. Moment matching methods

Next, we describe a variance reduction technique proposed by Barraquand (1995), who termed it quadratic resampling. His technique is based on moment matching. As before, we introduce it with the simple example of estimating the European call option price on a single asset and then generalize.

Let \( Z_i, i = 1, \ldots, n \), denote independent standard normals used to drive a simulation. The sample moments of the \( n \) \( Z \)'s will not exactly match those of the standard normal. The idea of moment matching is to transform the \( Z \)'s to match a finite number of the moments of the underlying population.
For example, the first moment of the standard normal can be matched by defining

$$\bar{Z}_i = Z_i - \bar{Z}, \quad i = 1, \ldots, n,$$

(8)

where $\bar{Z} = \frac{1}{n} \sum_{i=1}^{n} Z_i/n$ is the sample mean of the $Z_i$'s. Note that the $\bar{Z}_i$'s are normally distributed if the $Z_i$'s are normal. However, the $\bar{Z}_i$'s are not independent. As before, terminal stock prices are generated from the formula

$$S_T(i) = S_0 e^{(r - \sigma^2/2)T + \sigma \sqrt{T} \epsilon_i}, \quad i = 1, \ldots, n.$$

A moment-matched estimator of the call option price is the average of the $n$ values $C_i = e^{-rT} \max(S_T(i) - K, 0)$.

In the standard Monte Carlo method, confidence intervals for the true value $C$ could be estimated from the sample mean and variance of estimator. This cannot be done here since the $n$ values of $\bar{Z}$ are no longer independent, and hence the values $\bar{C}_i$ are not independent. This points out one drawback of the moment matching method: confidence intervals are not as easy to obtain. Indeed, for confidence intervals it appears to be necessary to apply moment matching to independent batches of runs and estimate the standard error from the batch means. This reduces the efficacy of the method compared with matching moments across all runs.

Eq. (8) showed one way to match the first moment of a distribution with mean zero. If the underlying population does not have a zero mean, transformed $Z$'s could be generated using $\bar{Z}_i = Z_i - \bar{Z} + \mu_Z$, where $\mu_Z$ is the population mean. The idea can easily be extended to match two moments of a distribution. In this case, an appropriate transformation is

$$\bar{Z}_i = (Z_i - \bar{Z}) \frac{\sigma_Z}{s_Z} + \mu_Z, \quad i = 1, \ldots, n,$$

(9)

where $s_Z$ is the sample standard deviation of the $Z_i$'s and $\sigma_Z$ is the population standard deviation. Of course, for a standard normal, $\mu_Z = 0$ and $\sigma_Z = 1$. An estimator of the call option price is the average of the $n$ values $\bar{C}_i$.

Using the transformation (9), the $\bar{Z}_i$'s are not normally distributed even if the $Z_i$'s are normal. Hence, the corresponding $\bar{C}_i$ are biased estimators of the true option value. For most financial problems of practical interest, this bias is likely to be small. However, the bias can be arbitrarily large in extreme circumstances.

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1 This point is not merely a minor technical issue. The sample variance of the $\bar{C}_i$'s is usually a poor estimate of $\text{Var}[\bar{C}_i]$. 
(even when only the first moment of the distribution is matched).\footnote{8} The dependence and bias in the moment matching method makes it difficult to quantify the improvement in general analytical terms.

The moment matching method is another example of the idea to ‘use what you know’. In this simple European option example, the mean and variance of the terminal stock price $S_T$ is also known. So the moment matching idea could be applied to the simulated terminal stock values $S_T(i)$. In this case, to match the first moment, define

$$\tilde{S}_T(i) = S_T(i) - \tilde{S}_T + \mu_{S_T},$$

where $\mu_{S_T} = S_0 e^{rT}$ and $\tilde{S}_T$ is the sample mean of the $S_T(i)$’s. To match the first two moments, define

$$\tilde{S}_T(i) = (S_T(i) - \tilde{S}_T) \frac{\sigma_{S_T}}{S_{S_T}} + \mu_{S_T},$$

where $\sigma_{S_T} = S_0 \sqrt{e^{2rT} (e^{\sigma^2 T} - 1)}$ and $S_{S_T}$ is the sample standard deviation of the $S_T(i)$’s. Duan and Simonato (1995) use a related method. They apply a multiplicative transformation to asset prices to enforce the martingale property over a finite set of paths.\footnote{9} They apply their method to GARCH option pricing.

Comparisons of various moment matching strategies are given in Table 1. For this comparison, $n = 100$ simulation trials were used to estimate the European call option price. Standard errors were estimated by re-simulation. That is, $m = 10,000$ simulation trials were conducted, each one based on $n$ replications of the estimator. The sample standard deviation of the $m$ simulation estimates gives an estimate of the standard error of a single simulation estimate. Root-mean-squared errors are not reported because they are identical to the standard errors for the number of digits reported.

The results in Table 1 show that matching two moments can reduce the simulation error by a factor ranging from 2 to 10. Matching two moments dominates matching one moment, but there is not a clear choice between transforming the original standard normals using (9) or the terminal stock prices

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\footnote{8}{For example, let $Z$ take the values $+1$ or $-1$ with probability one-half. Consider a security which pays $+1$ if $Z = 1$ and $-x$ if $Z \neq 1$. The expected payoff of the security is $(1 - x)/2$. To estimate this expected payoff by Monte Carlo simulation, draw $n$ samples $Z_i$ according to the prescribed distribution. Then use Eq. (8) to define $\tilde{Z}_i$ which match the first moment. For almost all samples for any large $n$, the estimated expected payoff is $-x$ and the bias is $(1 - x)/2$. This bias does not decrease as $n$ increases. Care must be taken when using Eqs. (8) or (9) when the support of the random variable is not the entire real line. For example, applying (8) or (9) to uniform or exponential random variables could cause the transformed values to fall outside of the relevant domain.}

\footnote{9}{This is equivalent to enforcing put-call parity.}
Table 1
Standard errors for European call option

<table>
<thead>
<tr>
<th>$\sigma$</th>
<th>$S_0/K$</th>
<th>No variance reduction</th>
<th>MM1 Eq. (8)</th>
<th>MM2 Eq. (9)</th>
<th>MM1 Eq. (10)</th>
<th>MM2 Eq. (11)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.2</td>
<td>0.9</td>
<td>0.24</td>
<td>0.19</td>
<td>0.11</td>
<td>0.19</td>
<td>0.09</td>
</tr>
<tr>
<td></td>
<td>1.0</td>
<td>0.62</td>
<td>0.29</td>
<td>0.09</td>
<td>0.26</td>
<td>0.10</td>
</tr>
<tr>
<td></td>
<td>1.1</td>
<td>0.93</td>
<td>0.19</td>
<td>0.09</td>
<td>0.15</td>
<td>0.11</td>
</tr>
<tr>
<td>0.4</td>
<td>0.9</td>
<td>0.80</td>
<td>0.55</td>
<td>0.24</td>
<td>0.51</td>
<td>0.17</td>
</tr>
<tr>
<td></td>
<td>1.0</td>
<td>1.22</td>
<td>0.66</td>
<td>0.19</td>
<td>0.56</td>
<td>0.23</td>
</tr>
<tr>
<td></td>
<td>1.1</td>
<td>1.61</td>
<td>0.63</td>
<td>0.17</td>
<td>0.48</td>
<td>0.28</td>
</tr>
<tr>
<td>0.6</td>
<td>0.9</td>
<td>1.40</td>
<td>0.95</td>
<td>0.38</td>
<td>0.84</td>
<td>0.28</td>
</tr>
<tr>
<td></td>
<td>1.0</td>
<td>1.93</td>
<td>1.10</td>
<td>0.31</td>
<td>0.91</td>
<td>0.39</td>
</tr>
<tr>
<td></td>
<td>1.1</td>
<td>2.38</td>
<td>1.13</td>
<td>0.25</td>
<td>0.85</td>
<td>0.49</td>
</tr>
</tbody>
</table>

All results are based on $n = 100$ simulation trials. The option parameters are: $K = 100$, $r = 0.10$, $T = 0.2$, with $S_0$ and $\sigma$ varying as indicated. Standard error estimates are based on $m = 10,000$ simulations.

using (11). Further computational results, not included in Table 1, indicate that the improvement factor with moment matching is essentially constant as $n$ increases. This may seem counterintuitive, since the moment matching adjustments converge to zero as $n$ increases. But the progressively smaller adjustments are equally important in reducing the estimation error as the number of simulation trials increases. For example, the standard error for $n = 10,000$ simulation trials is one-tenth of the corresponding number for $n = 100$ reported in Table 1.

The moment matching method can be extended to match covariances. For options that depend on multiple assets, the entire covariance structure is typically a simulation input. Barraquand (1995) suggests a method to match the entire covariance structure and reports error reduction factors ranging from two to several hundred for this method applied to pricing options on the maximum of $k$ assets.

The moment matching procedure could be applied to matching higher order moments as well. In addition to different methods for transforming random outcomes to match specified moments, additional points could be added as another way to match moments.

Whenever a moment is known, it can be used as a control rather than for moment matching. In an appendix, we give a theoretical argument favoring the use of moments as controls rather than for matching.

2.5. Stratified and Latin hypercube sampling

Like many variance reduction techniques, stratified sampling seeks to make the inputs to simulation more regular than random inputs. In particular, it
forces certain empirical probabilities to match theoretical probabilities, just as
moment matching forces empirical moments to match theoretical moments.

Consider, for example, the generation of 100 normal random variates as
inputs to a simulation. The empirical distribution of an independent sample
\( Z_1, \ldots, Z_{100} \) will look only roughly like the normal density; the tails of the
distribution—often the most important part—will inevitably be underrep-
resented. Stratified sampling can be used to force exactly one observation to lie
between the \((i-1)\)th and \(i\)th percentile, \( i = 1, \ldots, 100 \), and thus produce a better
match to the normal distribution. One way to implement this generates 100
independent random variates \( U_1, \ldots, U_{100} \), uniform on \([0, 1]\) and sets
\( \tilde{Z}_i = N^{-1}(\{i + U_i - 1\}/100), \quad i = 1, \ldots, 100, \) where \( N^{-1} \) is the inverse of the
cumulative normal distribution. This works because \((i + U_i - 1)/100\) falls be-
tween the \((i-1)\)th and \(i\)th percentiles of the uniform distribution, and percentiles
are preserved by the inverse transform.

Of course, \( \tilde{Z}_1, \ldots, \tilde{Z}_{100} \) are highly dependent, complicating the estimation of
standard errors. Computing confidence intervals with stratified sampling typically
requires batching the runs. For example, with a budget of 100,000 replications
we might run 100 independent stratified samples each of size 1000, rather
than a single stratified sample of size 100,000. To estimate standard errors we
must therefore sacrifice some variance reduction, just as with moment matching.

In principle, this approach applies in arbitrary dimensions. To generate a
stratified sample from the \(d\)-dimensional unit hypercube, with \(n\) strata in each
coordinate, we could generate a sequence of vectors
\[ U_j = (U_{j1}^{(1)}, \ldots, U_{jd}^{(d)}), \quad j = 1, 2, \ldots, \]
and then set
\[ V_j = \frac{U_j + (i_1, \ldots, i_d)}{n}, \quad i_k = 0, \ldots, n - 1, \quad k = 1, \ldots, d. \]

Exactly one \(V_j\) will lie in each of the \(n^d\) cubes defined by the product of the \(n\)
strata in each coordinate.

The difficulty in high dimensions is that generating even a single stratified
sample of size \(n^d\) may be prohibitive unless \(n\) is very small. Latin hypercube
sampling can be viewed as a way of randomly sampling \(n\) points of a stratified
sample while preserving some of the regularity from stratification. The method
was introduced by McKay et al. (1979) and further analyzed in Stein (1987). It
works as follows. Let \(\pi_1, \ldots, \pi_d\) be independent random permutations of
\(\{1, \ldots, n\}\), each uniformly distributed over all \(n!\) possible permutations. Set
\[ V_j^{(k)} = \frac{U_j^{(k)} + \pi_k(j) - 1}{n}, \quad k = 1, \ldots, d, \quad j = 1, \ldots, n. \]

The randomization ensures that each vector \(V_j\) is uniformly distributed over
the \(d\)-dimensional hypercube. At the same time, the coordinates are perfectly
stratified in the sense that exactly one of \( V_1^{(k)} \), \( \ldots \), \( V_n^{(k)} \) falls between \((j-1)/n\) and \( j/n, \ j = 1, \ldots, n\), for each dimension \( k = 1, \ldots, d\). As before, the dependence introduced by this method implies that standard errors can be estimated only through batching.

These methods can be viewed as part of a hierarchy of methods introducing additional levels of regularity in inputs at the expense of complicating the estimation of errors. Some, like stratified sampling, fix the size of the sample while others leave flexibility. The extremes of this hierarchy are straightforward Monte Carlo (completely random) and the low-discrepancy methods (completely deterministic) discussed in Section 3. Owen (1994, 1995) discusses these and other methods and introduces a hybrid that combines the regularity of low-discrepancy methods with the simple error estimation of standard Monte Carlo. Shaw (1995) uses an extension proposed by Stein (1987) to handle dependent inputs in a novel approach to estimating value at risk.

2.6. Some numerical comparisons

The variance reduction methods discussed thus far are fairly generic, in the sense that they do not rely on the detailed structure of the security to be priced. This contrasts with the remaining two methods that we discuss — importance sampling and conditional Monte Carlo. These methods must be carefully tailored to each application. It therefore seems appropriate to digress briefly into a numerical comparison of the generic methods on some option pricing problems.

We first examine the performance of these methods in pricing Asian options. The payoff of a discretely sampled arithmetic average Asian option is

\[
\max(S - K, 0),
\]

where \( S = \sum_{i=1}^{k} S_t / k \), \( S_t \) is the asset price at time \( t_i = iT/k \), and \( T \) is the option maturity. The value of the option is \( \mathbb{E}[e^{-rT}\max(S - K, 0)] \). There is no easily evaluated closed-form expression for this option value. Various formulas to approximate the Asian option price have been developed, but simulation is usually used to test the accuracy of the approximations.

For this Asian option, \( k \) random numbers are needed to simulate one option payoff, and \( nk \) random numbers are needed in total. Moment matching (MM2, for two moments) was applied \( k \) times to the \( n \) numbers used to generate each \( S_t \) at time \( t_i \). Latin hypercube sampling (LHS) was applied to sample \( n \) points from the \( k \)-dimensional unit cube. The discretely sampled geometric average Asian option price was used as a control variate (see Turnbull and Wakeman (1991) for a closed-form solution for this price). Results appear in Table 2.

The results in Table 2 indicate that matching two moments can reduce the simulation error by a factor ranging from 1 to 10. Using the geometric average Asian option price as a control variate reduces error by a factor ranging from 20
Table 2
Standard errors for arithmetic average Asian options

<table>
<thead>
<tr>
<th>$\sigma$</th>
<th>$K/S_0$</th>
<th>No variance reduction</th>
<th>Antithetic method</th>
<th>Control variate</th>
<th>MM2</th>
<th>LHS</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.2</td>
<td>0.9</td>
<td>0.053</td>
<td>0.052</td>
<td>0.003</td>
<td>0.048</td>
<td>0.049</td>
</tr>
<tr>
<td></td>
<td>1.0</td>
<td>0.344</td>
<td>0.231</td>
<td>0.004</td>
<td>0.162</td>
<td>0.161</td>
</tr>
<tr>
<td></td>
<td>1.1</td>
<td>0.566</td>
<td>0.068</td>
<td>0.006</td>
<td>0.052</td>
<td>0.058</td>
</tr>
<tr>
<td>0.4</td>
<td>0.9</td>
<td>0.308</td>
<td>0.297</td>
<td>0.014</td>
<td>0.240</td>
<td>0.248</td>
</tr>
<tr>
<td></td>
<td>1.0</td>
<td>0.694</td>
<td>0.506</td>
<td>0.017</td>
<td>0.352</td>
<td>0.354</td>
</tr>
<tr>
<td></td>
<td>1.1</td>
<td>1.017</td>
<td>0.388</td>
<td>0.021</td>
<td>0.281</td>
<td>0.289</td>
</tr>
<tr>
<td>0.6</td>
<td>0.9</td>
<td>0.632</td>
<td>0.583</td>
<td>0.032</td>
<td>0.451</td>
<td>0.455</td>
</tr>
<tr>
<td></td>
<td>1.0</td>
<td>1.052</td>
<td>0.817</td>
<td>0.038</td>
<td>0.566</td>
<td>0.578</td>
</tr>
<tr>
<td></td>
<td>1.1</td>
<td>1.443</td>
<td>0.759</td>
<td>0.047</td>
<td>0.539</td>
<td>0.560</td>
</tr>
</tbody>
</table>

All results are based on $n = 100$ simulation trials with $k = 50$ prices in the average. The option parameters are: $K = 100$, $r = 0.10$, $T = 0.2$, with $S_0$ and $\sigma$ varying as indicated. Standard error estimates based on $m = 10,000$ simulations. The geometric average Asian option is used as the control variate. Moment matching (MM2) was applied to the $i$th price in the average, $i = 1, \ldots$, across replications.

to 100, and is consistently the most effective method. LHS and MM2 perform similarly. Antithetics are consistently dominated by the other methods.

Next, we compare these variance reduction techniques in pricing down-and-out call options with discrete barriers. The payoff of this option at expiration is the standard call option payoff if the asset price $S_t$ exceeds the barrier $H$ at all times $t_i = iT/k$, $i = 1, \ldots, k$, otherwise the payoff is zero. The option is knocked out if $S_t \leq H$ at any time $t_i$. As a control we use the Black-Scholes price of a standard call. Moment matching and LHS are implemented as with the Asian option. Results are given in Table 3. These are consistent with the pattern in Table 2, except that the superiority of the control variate method is less pronounced.

Although it is always risky to draw conclusions from limited numerical evidence, we suggest the following broad conclusions. The antithetic method is easy to implement, but often leads to only modest error reductions. Moment matching is similarly easy to implement and often leads to significant error reductions, but the error estimation is more difficult and bias is a potential problem. LHS suffers from the same error estimation difficulty but does not introduce bias. The control variate technique can lead to very substantial error reductions, but its effectiveness hinges on finding a good control for each problem.
Table 3
Standard errors for down-and-out call options with discrete barriers

<table>
<thead>
<tr>
<th>( \sigma )</th>
<th>( K/S_0 )</th>
<th>No variance reduction</th>
<th>Antithetic method</th>
<th>Control variate</th>
<th>MM2</th>
<th>LHS</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.2</td>
<td>0.9</td>
<td>0.96</td>
<td>0.44</td>
<td>0.37</td>
<td>0.43</td>
<td>0.39</td>
</tr>
<tr>
<td>1.0</td>
<td>0.9</td>
<td>0.62</td>
<td>0.44</td>
<td>0.13</td>
<td>0.31</td>
<td>0.30</td>
</tr>
<tr>
<td>1.1</td>
<td>0.8</td>
<td>0.30</td>
<td>0.28</td>
<td>0.03</td>
<td>0.22</td>
<td>0.22</td>
</tr>
<tr>
<td>0.2</td>
<td>0.9</td>
<td>1.59</td>
<td>1.15</td>
<td>0.73</td>
<td>0.95</td>
<td>0.88</td>
</tr>
<tr>
<td>1.0</td>
<td>1.22</td>
<td>1.00</td>
<td>0.45</td>
<td>0.76</td>
<td>0.74</td>
<td></td>
</tr>
<tr>
<td>1.1</td>
<td>0.88</td>
<td>0.82</td>
<td>0.26</td>
<td>0.61</td>
<td>0.61</td>
<td></td>
</tr>
<tr>
<td>0.2</td>
<td>0.9</td>
<td>2.19</td>
<td>1.83</td>
<td>1.07</td>
<td>1.44</td>
<td>1.36</td>
</tr>
<tr>
<td>1.0</td>
<td>1.86</td>
<td>1.62</td>
<td>0.80</td>
<td>1.25</td>
<td>1.23</td>
<td></td>
</tr>
<tr>
<td>1.1</td>
<td>1.54</td>
<td>1.40</td>
<td>0.58</td>
<td>1.09</td>
<td>1.09</td>
<td></td>
</tr>
</tbody>
</table>

All results are based on \( n = 100 \) simulation trials. There are \( k = 5 \) points in the discrete barrier at 95. The other option parameters are: \( S_0 = 100, r = 0.10, T = 0.2 \), with \( K \) and \( \sigma \) varying as indicated. Standard error estimates are based on \( m = 10,000 \) simulations. The standard European call option (Black-Scholes formula) is used as the control variate. Moment matching (MM2) was applied to the \( i \)th return, \( i = 1, \ldots, 5 \), across replications.

2.7. Importance sampling

This technique builds on the observation that an expectation under one probability measure can be expressed as an expectation under another through the use of a likelihood ratio or Radon-Nikodym derivative. This idea is familiar in finance because it underlies the representation of prices as expectations under a martingale measure. In Monte Carlo, the change of measure is used to try to obtain a more efficient estimator. We present some examples using this technique; for general background see Bratley et al. (1987) or Hammersley and Handscomb (1964).

As a simple example, consider the evaluation of the Black-Scholes price of a call option, i.e., the computation of \( e^{-rT} \mathbb{E} \{ \max \{ S_T - K, 0 \} \} \) with \( S_T \) as in (2). A straightforward approach generates samples of the terminal value \( S_T \) consistent with a geometric Brownian motion having drift \( r \) and volatility \( \sigma \), just as in (2). But we are in fact free to generate \( S_T \) consistent with any other drift \( \mu \), provided we weight the result with a likelihood ratio. For emphasis, we subscript the expectation operator with the drift parameter. Then

\[
E_{r} \{ \max \{ S_T - K, 0 \} \} = E_{\mu} \{ \max \{ S_T - K, 0 \} L \}.
\]
where the likelihood ratio $L$ is the ratio of the log-normal densities with parameters $r$ and $\mu$ evaluated at $S_T$, given by

$$L = \left( \frac{S_T}{S_0} \right)^{r - \mu/s^2} \exp \left( \frac{(\mu^2 - r^2)T}{2s^2} \right).$$

Indeed, $S_T$ need not even be sampled from a log-normal distribution. The only requirement is that the support of the importance sampling measure contain the support of the original measure so that the likelihood ratio is well-defined; this is an absolute continuity requirement. In the example above, this means that any distribution for $S_T$ whose support includes $(0, \infty)$ is admissible.

Ideally, one would like to choose the importance sampling distribution to reduce variance. In the example above, one obtains a zero-variance estimator by sampling $S_T$ from the density

$$f(x) = c^{-1} \max \{x - K, 0\} e^{-rT} g(x),$$

where $g$ is the (log-normal) density of $S_T$ and $c$ is a normalizing constant that makes $f$ integrate to 1. The difficulty is that $c$ is the Black-Scholes price itself, so this method requires knowledge of the solution for its implementation. Nevertheless, it gives some indication of the potential gain from importance sampling.

Reider (1993) has investigated the impact of importance sampling based on a change of drift and volatility. (Changing the volatility is consistent with absolute continuity in a discrete-time approximation of a diffusion though not in the continuous-time limit.) He finds that choosing the importance sampling distribution to have higher drift and volatility provides substantial variance reduction in pricing deep out-of-the-money options. He also investigates the combination of importance sampling with antithetic variates and control variates, and the use of put-call parity for indirect estimation. Nielsen (1994) has explored some related importance sampling ideas in sampling from a binomial tree.

Andersen (1995) has developed a powerful application of importance sampling for simulating interest rates and has applied it to nonlinear stochastic differential equation models. We briefly describe his approach. Let $r_t$ be the instantaneous short rate described, e.g., by a diffusion model. Then

$$B(T) = E \left[ \exp \left( - \int_0^T r_t \, dt \right) \right]$$
is the price today of a zero-coupon bond with face value $1$, maturing at time $T$. In, for example, the Cox–Ingersoll–Ross and Vasicek models, $B(T)$ is available in closed form. We may therefore define a new probability measure $\tilde{P}$ by setting

$$\tilde{P}(A) = E \left[ \exp \left( - \int_0^T r_t \, dt - \log B(T) \right) \mathbf{1}_A \right]$$

for any event $A$, where $\mathbf{1}_A$ denotes the indicator of the event $A$. Let $E$ denote expectation with respect to $\tilde{P}$. Then for any random variable $X$, $E[X] = \tilde{E}[X \mid L_T]$ where the likelihood ratio $L_T$ is given by

$$L_T = \exp \left( \int_0^T r_t \, dt + \log B(T) \right).$$

In particular, if we take $X = \exp \left( - \int_0^T r_t \, dt \right)$, we know that $E[X] = B(T)$ and, therefore, $B(T)$ is the expectation under $\tilde{P}$ of $X \mid L_T$; i.e., of

$$\exp \left( - \int_0^T r_t \, dt \right) \times \exp \left( \int_0^T r_t \, dt + \log B(T) \right).$$

But this simplifies to $B(T)$ itself, meaning that we obtain a zero-variance estimator of the bond price by switching to the new probability measure. Moreover, Andersen shows that sample paths of $r_t$ can be generated under $\tilde{P}$ simply by applying a change of drift to the original process.

As described above, the method would appear to require knowledge of the solution for its implementation. Nevertheless, the method has two important applications. The first is in the pricing of contingent claims. Because $\tilde{P}$ eliminates the variance of bond prices, it should be effective in reducing variance for pricing, e.g., European bond options expiring at time $T$. Andersen's numerical results bear this out. A second application is in the pricing of bond models with no closed-form solutions: Andersen's results show that the change of drift derived from a tractable model (like CIR or Vasicek) remains effective when applied to an intractable model, and this significantly expands the scope of the method.

Importance sampling is frequently used to make rare events less rare; this is already suggested in Reider's (1994) application to out-of-the-money options. Our next example further highlights this aspect through a new application to barrier options. We consider a knock-in option far from the barrier and use importance sampling to increasing the probability of a payout.

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10See, e.g., Hull (1997, Chapter 17) for background on these models.
Suppose the barrier is monitored at discrete times $n \Delta t$, $n = 0, 1, \ldots, m$, with $\Delta T = T/m$. Set the barrier at $H = S_0 e^{-b}$ and the strike at $K = S_0 e^c$, with $b, c > 0$. A down-and-in call pays $S_T - K$ at time $T$ if $S_T > K$ and $S_{n \Delta t} < H$ for some $n = 1, \ldots, m$. We can write the price of the underlying at monitoring instants as

$$S_{n \Delta t} = S_0 e^{u_i}, \quad U_n = \sum_{i=1}^{n} X_i,$$

with the $X_i$ i.i.d. normal having mean $(r - \frac{1}{2} \sigma^2) \Delta t$ and variance $\sigma^2 \Delta t$. Let $\tau$ be the first time $U_n$ drops below $-b$; then the probability of a payout is $P(\tau < m, U_m > c)$. If $b$ and $c$ are large, this probability is small, and most simulation runs return zero. Through importance sampling, we can increase this probability and thus get more information out of each run.

Consider alternative probability measures $P_{\mu_1, \mu_2}$ that give $U_n$ a drift of $\mu_1 \Delta t$ until $\tau$ and then switch the drift to $\mu_2 \Delta t$. Intuitively, we would like to make $\mu_1 < 0$ to drive the asset price to the barrier and then make $\mu_2 > 0$ to drive it above the strike. For any $\mu_1, \mu_2$, we have

$$P(\tau < m, U_m > c) = E_{\mu_1, \mu_2} \{ L_{\mu_1, \mu_2} I_{[\tau < m, U_m > c]} \}.$$

The likelihood ratio is given by

$$L_{\mu_1, \mu_2} = \exp(-\theta_1 U_\tau + \psi(\theta_1) \tau - \theta_2 (U_m - U_\tau) + \psi(\theta_2)(m - \tau)),$$

where $\theta_i = (\mu_i - r + \frac{1}{2} \sigma^2)/\sigma^2$, $i = 1, 2$, and $\psi(\theta) = (r - \frac{1}{2} \sigma^2) \Delta t \theta + \frac{1}{2} \sigma^2 \Delta t \theta^2$. This follows from algebraic simplification of the product of the ratios of the densities of the $X_i$ under the original and new means.

It remains to choose $\mu_1, \mu_2$. Intuitively, most of the variability in $L_{\mu_1, \mu_2}$ comes from $\tau$ (the time of the barrier crossing); for large $b, c$, in the event of a payout we expect to have $U_\tau \approx -b$ and $U_m \approx c$ so these terms should contribute less variability. If we choose $\mu_1, \mu_2$ so that $\psi(\theta_1) = \psi(\theta_2)$, the likelihood ratio simplifies to

$$L_{\mu_1, \mu_2} = \exp(-(\theta_1 - \theta_2)U_\tau - \theta_2 U_m + m\psi(\theta_2)),$$

which depends on $\tau$ only through $U_\tau \approx -b$. The condition $\psi(\theta_1) = \psi(\theta_2)$ translates to $\mu_1 = -\mu_2 \equiv -\mu$, so it only remains to choose this drift parameter. We choose it so that the time to traverse the straight line path from 0 to $-b$ and then to $c$ at rate $\mu$ equals the number of steps $m$:

$$\frac{b}{\mu \Delta t} + \frac{(b + c)}{\mu \Delta t} = m;$$
Table 4
Standard errors for down-and-in calls: importance sampling

<table>
<thead>
<tr>
<th>$H$</th>
<th>$K$</th>
<th>No variance reduction</th>
<th>Importance sampling</th>
<th>Efficiency ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>92</td>
<td>100</td>
<td>0.00309</td>
<td>0.00069</td>
<td>20</td>
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<tr>
<td>92</td>
<td>105</td>
<td>0.00129</td>
<td>0.00014</td>
<td>85</td>
</tr>
<tr>
<td>88</td>
<td>96</td>
<td>0.00110</td>
<td>0.00011</td>
<td>96</td>
</tr>
<tr>
<td>85</td>
<td>90</td>
<td>0.00084</td>
<td>0.00008</td>
<td>116</td>
</tr>
<tr>
<td>92</td>
<td>105</td>
<td>0.01418</td>
<td>0.00541</td>
<td>7</td>
</tr>
<tr>
<td>85</td>
<td>105</td>
<td>0.00328</td>
<td>0.00038</td>
<td>75</td>
</tr>
<tr>
<td>75</td>
<td>96</td>
<td>0.00030</td>
<td>0.00001</td>
<td>1124</td>
</tr>
<tr>
<td>75</td>
<td>85</td>
<td>0.00148</td>
<td>0.00010</td>
<td>222</td>
</tr>
</tbody>
</table>

All results are based on $n = 100,000$ simulation trials. The parameters are: $S_0 = 95$, $r = 0.15$, and $r = 0.05$, with the barrier $H$ and strike $K$ varying as indicated. The first four cases have $T = 0.25$ and $m = 50$; the last four have $T = 1$ and $m = 250$.

i.e., $\mu = (2b + c)/T$. Interestingly, this change of drift does not depend on the original mean increment $(r - \frac{1}{2} \sigma^2)At$.

Table 4 illustrates the performance of this method. The computational effort with and without importance sampling is essentially the same, so the efficiency improvement is just the ratio of the variances. The improvement varies widely but shows the potential for dramatic gains from importance sampling, particularly when the barrier is far from the current price of the underlying.\textsuperscript{11}

In recent work, Andersen and Brotherton-Ratcliffe (1996) and Beaglehole, Dybuig, and Zhou (1997) show how to eliminate the bias caused by using a simulation at a discrete set of times to price continuous options on extrema, e.g., barrier or lookback options.

2.8. Conditional Monte Carlo

This approach to efficiency improvement exploits the variance reducing property of conditional expectation: for any random variables $X$ and $Y$, $\text{Var} [E[X \mid Y]] \leq \text{Var} [X]$, with strict inequality except in trivial cases.\textsuperscript{12} In replacing an estimator by its conditional expectation we reduce variance essentially because we are doing part of the integration analytically and leaving less to be done by Monte Carlo.

\textsuperscript{11} The standard errors in the table are all quite small, but so are the associated option values. Hence, the relative error without importance sampling is quite significant.

\textsuperscript{12} This is a direct consequence of Jensen's inequality for conditional expectations.
Hull and White (1987) use this idea to price options with stochastic volatilities. Consider a model in which an asset price and its variability evolve as follows:

$$dS = rS dt + \nu S dW_1, \quad dv^2 = zv^2 dt + \xi v^2 dW_2,$$

with $W_1, W_2$ independent. Suppose we want to price a standard European call on $S$. A straightforward approach simulates sample paths of $v$ and $S$ up to time $T$ and averages $\max\{S_T - K, 0\}$ over all paths. An alternative notes that, conditional on the path of $v_t$ in $[0, T]$, the asset price $S_t$ may be treated as having a time varying but deterministic volatility. Thus, conditional on the volatility path, the option can be priced by the Black-Scholes formula:

$$e^{-rt} E[\max\{S_T - K, 0\} | v_t, 0 \leq t \leq T] = \text{BS}(S_0, K, r, T, \sqrt{V_T}),$$

where

$$V_T = \frac{1}{T} \int_0^T v_t^2 dt$$

is the average squared volatility over the path, and BS$(S, K, T, r, \sigma)$ is the Black-Scholes price of a call with constant volatility $\sigma$ and the other parameters as indicated. Using this conditional expectation as the estimator is sure to reduce variance and may even reduce computational effort since it obviates simulation of $S$. It is worth emphasizing that both straightforward Monte Carlo and conditional Monte Carlo would have to be applied to discrete-time approximations of the continuous processes above. Also, the applicability of conditional Monte Carlo in this setting relies critically on the fact that the evolution of the asset price does not influence the volatility path. See Willard (1996) for an extension to the case of correlated $W_1$ and $W_2$.

As a further illustration of the use of conditional Monte Carlo, we give a new illustration in the pricing of a down-and-in call with a discretely monitored barrier. Let $0 = t_0 < t_1 < \cdots < t_n = T$ be the monitoring instants and $S_t$ the price of the underlying at the $i$th such instant. The option price is $E[e^{-rt} \max\{S_T - K, 0\} 1_{t_n \leq T}]$, where $H$ is the barrier and $\tau_H$ is the first monitoring time at which the barrier is breached.

Straightforward simulation generates paths of the underlying and evaluates the estimator

$$e^{-rt} \max\{S_T - K, 0\} 1_{\tau_H \leq T},$$

Our first alternative conditions on $\{S_0, \ldots, S_n\}$, the path of the underlying until the barrier crossing; i.e.,

$$E[e^{-rt} \max\{S_T - K, 0\} 1_{\tau_H \leq T}]$$

$$= e^{-rt} E[\max\{S_T - K, 0\} 1_{\tau_H \leq T} | S_0, \ldots, S_n]$$

$$= e^{-rt} E[\text{BS}(S_{\tau_H}, K, r, T - \tau_H, \sigma) 1_{\tau_H \leq T}].$$
This yields the estimator

$$
CMC_1 = e^{-rT} BS(S_{t_n}, K, r, T - \tau_H, \sigma) 1_{\{t_n \leq T\}}.
$$

This says: simulate until the barrier is crossed or the option expires; if the barrier was crossed, return the Black-Scholes price starting from price $S_{t_n}$ with maturity $T - \tau_H$.

Our second alternative conditions one step earlier, at each monitoring instant evaluating the probability that the barrier will be breached for the first time at the next monitoring instant:

$$
E[e^{-rT} \max\{S_T - K, 0\} 1_{\{t_n \leq T\}}]
$$

$$
= e^{-rT} E\left[ \max\{S_T - K, 0\} \sum_{n=1}^{m} 1_{\{t_n - t_n\}} \right]
$$

$$
= e^{-rT} E\left[ \sum_{n=1}^{m} E[\max\{S_T - K, 0\} 1_{\{t_n = t_n\}} | S_{t_n}, \ldots, S_{t_n-1}] \right]
$$

$$
= e^{-rT} E\left[ \sum_{n=0}^{t_n-1} BS2(S_{t_n}, K, H, r, t_{n+1} - t_n, T - t_n, \sigma) \right]
$$

where $BS2(S, K, H, r, t, T, \sigma)$ is the price of a down-and-in call that knocks in only if the underlying is below $H$ at time $t$. We thus arrive at the estimator

$$
CMC_2 = e^{-rT} \sum_{n=0}^{t_n-1} BS2(S_{t_n}, K, H, r, t_{n+1} - t_n, T - t_n, \sigma),
$$

with

$$
BS2(S, K, H, r, t, T, \sigma) = SN_2(a_1, b_1, \rho) - e^{-rT} KN_2(a_2, b_2, \rho),
$$

where $\rho = -\sqrt{t/T}$, $N_2$ is the bivariate cumulative normal distribution with correlation $\rho$, and

$$
a_1 = \frac{\log(S/K) + (r + \frac{1}{2} \sigma^2) T}{\sigma \sqrt{T}}, \quad a_2 = a_1 - \sigma \sqrt{T},
$$

$$
b_1 = \frac{\log(H/S) - (r + \frac{1}{2} \sigma^2) t}{\sigma \sqrt{t}}, \quad b_2 = b_1 + \sigma \sqrt{t}.$$
Table 5
Comparison of CMC estimators for down-and-in call

<table>
<thead>
<tr>
<th>Method</th>
<th>Standard error (s)</th>
<th>Computation time (t)</th>
<th>s√t</th>
</tr>
</thead>
<tbody>
<tr>
<td>Base</td>
<td>0.108</td>
<td>0.133</td>
<td>0.039</td>
</tr>
<tr>
<td>CMC₁</td>
<td>0.034</td>
<td>0.117</td>
<td>0.012</td>
</tr>
<tr>
<td>CMC₂</td>
<td>0.021</td>
<td>3.233</td>
<td>0.038</td>
</tr>
<tr>
<td>CMC</td>
<td>0.014</td>
<td>3.367</td>
<td>0.026</td>
</tr>
</tbody>
</table>

Results based on n = 10,000 replications with σ = 0.4, r = 0.10, S₀ = K = 100, H = 95, T = 0.5, and 10 equally spaced monitoring times.

(The derivation of this formula is fairly standard and therefore omitted.) The CMC₂ estimator can be expected to have lower variance than the CMC₁ estimator because it conditions on less information and thus does more integration analytically. In fact, CMC₂ is not a conditional Monte Carlo estimator in the strict sense because it conditions on different information at different times, making it more precisely a filtered Monte Carlo estimator in the sense of Glasserman (1996).

Because the two estimators above have the same expectation, their difference has mean 0 and can be used as a control variate to form a further estimator

\[ CMC' = CMC₁ + β(CMC₂ - CMC₁). \]

With β optimized, this has lower variance than either individual estimator.

Numerical results appear in Table 5. As expected, each level of conditioning further reduces variance, and the combined estimator achieves the lowest standard error of all. However, repeated evaluation of the function BS2 turns out to be time-consuming, making CMC₁ overall the most efficient estimator.

3. Low-discrepancy sequences

For complex problems the performance of the basic Monte Carlo approach may be rather unsatisfactory because the error is \( O(1/\sqrt{n}) \). We can sometimes improve convergence by using pre-selected deterministic points to evaluate the integral. The accuracy of this approach depends on the extent to which these deterministic points are evenly dispersed throughout the domain of integration. Discrepancy measures the extent to which the points are evenly dispersed throughout a region: the more evenly dispersed the points are the lower the discrepancy. Low-discrepancy sequences are often called quasi-random...
sequences even though they are not at all random.\textsuperscript{13} We shall use both terms in this paper.

Low-discrepancy methods have recently been used to tackle a number of problems in the finance area. These applications are more fully described in papers by Birge (1994), Joy et al. (1996) and Paskov and Traub (1995); the use of quasi-Monte Carlo is also proposed in Cheyette (1992). In this section we describe how the approach works and review some of the recent applications. The book by Press et al. (1992) provides an intuitive introduction to low-discrepancy sequences and quasi-Monte Carlo methods. Spanier and Maize (1994) provide a recent overview of quasi-random methods and how they can be used to evaluate integrals with medium sized samples. Niederreiter (1992) and Tezuka (1995) provide in-depth analyses of low-discrepancy sequences. Moskowitz and Caflisch discuss recent developments in improving the convergence of quasi-random Monte Carlo methods. In earlier work, Haselgrove (1961) describes a method for multivariate integration that can be applied to security pricing. Haselgrove’s method is developed for problems of eight dimensions or less and our numerical experiments suggest that it is competitive with the low-discrepancy sequences investigated in this section for problems of this size.

The basic idea behind the approach is quite intuitive and is readily explained in the one-dimensional case. Suppose we wish to integrate a function \( f(x) \) over the interval \([0, 1]\) using a sequence of \( n \) points. Rather than pick a random sequence suppose we pick a deterministic sequence of points that are, in some sense, evenly distributed. With this choice, the accuracy of the estimate will be higher than that obtained using the crude Monte Carlo approach. If we use an equally spaced grid we obtain the trapezoidal method of numerical integration which has an error of \( O(n^{-2}) \). However, the more challenging task is to evaluate multi-dimensional integrals. Without loss of generality, we can assume that the domain of integration is contained in the \( d \)-dimensional unit hypercube. The advantages of the uniformly spaced grid in the one-dimensional case do not carry over to higher dimensions. The principal reason is that the error bound for the \( d \)-dimensional trapezoidal rule is \( O(n^{-2/d}) \). In addition, if we use an evenly spaced Cartesian grid, we would have to decide the number of points in advance to achieve uniformity. This is restrictive because, in numerical applications, we would like to be able to add points sequentially until some termination criterion is met.

Low-discrepancy sequences have the property that as successive points are added the entire sequence of points still remains more or less evenly dispersed throughout the region. Niederreiter (1992) gives a detailed analysis of the discrepancy of a sequence. Here, we just briefly recall the definition. Suppose we

\textsuperscript{13} Thus, the name quasi-random is very misleading since these sequences are deterministic. However, it seems to be sanctioned by usage.
have a sequence of \( n \) points \( \{x_1, x_2, \ldots, x_n\} \) in the \( d \)-dimensional half-open unit cube, \( I^d = [0, 1)^d \) and a subset \( J \) of \( I^d \). We define

\[
D(J; n) = \frac{A(J; n)}{n} - V(J),
\]

where \( A(J; n) \) is the number of \( k, 1 \leq k \leq n \), with \( x_k \in J \) and \( V(J) \) is the volume of \( J \). The discrepancy, \( D_n \), of the sequence is defined to be the supremum of \( |D(J; n)| \) over all \( J \). The star discrepancy \( D^*_n \), is obtained by taking the supremum over sets \( J \) of the form

\[
\prod_{i=1}^d [0, u_i).
\]

In the one-dimensional case there is a simple explicit form for the (star)\(^{14}\) discrepancy of a sequence of \( n \) points. If we label the points so that, \( 0 \leq x_1 \leq \cdots \leq x_n \leq 1 \), then the discrepancy of this sequence is

\[
D^*_n = \frac{1}{2n} + \max_{k=1, \ldots, m} \left| x_k - \frac{2k - 1}{2n} \right|.
\]

We can see that the star discrepancy is at least \( 1/(2n) \) and that the lowest value is attained when

\[
x_k = \frac{2k - 1}{2n}, \quad 1 \leq k \leq n.
\]

In higher dimensions there is no simple form for the discrepancy of a sequence.

There are several examples of low-discrepancy sequences, including the sequences proposed by Halton (1960), Sobol' (1967), Faure (1982), and Niederreiter (1988).\(^{15}\) For these sequences the asymptotic form of the star discrepancy

\[^{14}\text{For the rest of the paper we simply use the term discrepancy rather than star discrepancy to refer to } D^*_n.\]

\[^{15}\text{Interestingly, linear congruential generators – frequently used to generate the pseudo-random numbers that drive ordinary Monte Carlo – produce sets of points with low-discrepancy over the entire period of the generator; see Niederreiter (1976). This suggests the possibility of choosing such a generator with period roughly equal to the total number of points required as a type of quasi-Monte Carlo method. In ordinary Monte Carlo, one prefers instead that the period be many orders of magnitude larger than the number of points required. We thank Peter Hellekalek of the University of Salzburg for this observation.}\]
has been shown to be

\[ D_n^* = O\left(\frac{\log n}{n}\right). \]

This bound for the discrepancy involves a constant which, in general, depends on the dimension \( d \) of the sequence. These constants are very difficult to estimate accurately in high dimensions. For large values of \( d \) the constants are often ridiculously large for reasonable values of \( n \) according to Spanier and Maize (1994, p. 23). Furthermore for high dimensions it may take a long time before the discrepancy reaches its asymptotic level. Morokoff and Caflisch (1995) note that for intermediate values of \( n \) the discrepancy may be \( O(\sqrt{n}) \). They suggest that the transition to \( O(n^{-1}(\log n)^d) \) occurs at around values of \( n = e^d \). For large \( d \) this will be an enormous number.

The error in numerical integration using a low-discrepancy sequence admits a deterministic bound. The bound reflects both the discrepancy of the sequence of points used to evaluate the integral as well as the regularity of the function. The result is contained in the following theorem.

**Theorem (Koksma–Hlawka).** Let \( I^d = [0, 1]^d \) and let \( f \) have bounded variation \( V(f) \) on \( [0, 1]^d \) in the Hardy-Krause\(^{16}\) sense. Then for any \( x_1, x_2, \ldots, x_n \in I^d \), we have

\[ \left| \frac{1}{n} \sum_{k=1}^{n} f(x_k) - \int_{I^d} f(u) \, du \right| \leq V(f)D_n^*. \]

The error bound provided by this theorem, while it is of theoretical interest, is of little help in most practical situations. The theoretical bound normally overestimates the actual error by a wide margin and \( V(f) \) may be difficult to evaluate or even approximate. We have noted that the constants buried in the bounds for the discrepancy are large. Another reason for the coarseness of the bound is that the Koksma–Hlawka theorem does not reflect additional smoothness in \( f \). Intuitively we would expect the approximation to be better as \( f \) becomes smoother. In finance applications the payoffs are normally continuous functions of the variables (with some important exceptions – payoffs on digital and barrier options are discontinuous), but may not be sufficiently smooth to have finite variation because of functions like ‘max’ embedded in the payoffs. Hlawka (1971) provides an alternative bound under weaker smoothness requirements.

\(^{16}\) For a more complete discussion of the Hardy-Krause definition of variation and details on this theorem see Niederreiter (1992).
To date, studies using low-discrepancy sequences in finance applications find that the errors produced are substantially lower than the corresponding errors generated by crude Monte Carlo. Joy et al. (1996) used Faure sequences to price several complex derivative securities. They found that the quasi-Monte Carlo approach resulted in significantly smaller errors than the standard Monte Carlo approach. They confirmed that the actual error bound (for cases in which it could be computed precisely) was dramatically less than the bound computed from the Klekma–Hlawka inequality. Paskov and Traub (1995) used both Sobol' sequences and Halton sequences to evaluate mortgage-backed security prices. Their work involves the evaluation of integrals with dimensions up to 360; they find that Sobol' sequences are more efficient than Halton sequences and that the quasi-random approach outperforms the standard Monte Carlo approach for these types of problems.17 Paskov and Traub's results stand in contrast to the claim that is sometimes found in the literature18 that the superiority of low-discrepancy algorithms vanishes for intermediate values of d around 30. Bratley et al. (1992) conducted practical numerical experiments using low-discrepancy sequences and conclude that standard Monte Carlo is superior to quasi-Monte Carlo for high dimensions, say greater than 12. They used Sobol' and Niederreiter sequences in their tests. They conclude that in high dimensions, 'quasi-Monte Carlo seems to offer no practical advantage over pseudo-Monte Carlo because the discrepancy bound for the former is far larger than \(\sqrt{n}\) for \(n = 2^{30}\), say'. (In a personal communication, Fox adds that the crossover probably depends a lot on the sequence.) The reason for the difference between this verdict and the results of the finance applications may be that the integrands typically found in finance applications behave better than those used by numerical analysts19 to compare different algorithms. Another important consideration is that financial applications typically involve discounting, and this may effectively reduce dimensionality; for example, some of the 360 months in the life of a mortgage may have little influence on the value of a mortgage-backed security. Nevertheless, the experience of Bratley et al. (1992) serves as a useful caution against assuming that quasi-Monte Carlo will outperform standard Monte Carlo in all situations.

17 Bratley et al. (1992) note that the Niederreiter sequence they tested theoretically beats Sobol's sequences in dimensions higher than seven.

18 See, for example, Rensburg and Torrie (1993) or Morokoff and Caflisch (1995).

19 For example, one of the integrals used by Bratley et al. (1992) was

\[
\int_0^1 \cdots \int_0^1 \sum_{k=1}^d k \cos(kx_t) \, dx_1 \cdots dx_d.
\]

This integrand is highly periodic for large values of \(d\).
Some theoretical differences among low-discrepancy sequences can be understood through the concepts of \((t, m, s)\)-nets and \((t, s)\)-sequences; these are discussed in detail in Niederreiter (1992). Briefly, an elementary interval in base \(b\) in dimension \(s\) is a set of the form
\[
\prod_{j=1}^{i} \left[ \frac{a_j}{b^{k_j}}, \frac{a_j + 1}{b^{k_j}} \right),
\]
with \(k_j, a_j\) nonnegative integers and \(a_j < b^{k_j}\). A \((t, m, s)\)-net (with \(0 \leq t \leq m\)) is a set of \(b^m\) points in the \(s\)-dimensional hypercube such that every elementary interval of volume \(b^{-m}\) contains \(b^t\) points. Speaking loosely, this means that the proportion of points in each sufficiently large box equals the volume of the box. Smaller \(t\) implies greater uniformity. An infinite sequence forms a \((t, s)\)-sequence if for all \(m \geq t\) certain finite subsequences of length \(b^m\) form \((t, m, s)\)-nets in base \(b\). Sobol' points are \((t, s)\)-sequences in base 2 and Faure points are \((0, s)\)-sequences in prime bases not less than \(s\). Thus, Faure points achieve the smallest value of \(t\), but at the expense of a large base. A smaller base implies that uniformity holds over shorter subsequences.

An important issue in the use of quasi-Monte Carlo concerns the termination criterion, since Koksma–Hlawka bound is often of little practical value. Various heuristics are available. Birge (1994) suggests that a rough bound may be obtained by tracking the maximum and minimum values over a period that shows equal numbers of increases and decreases. For instance, the criterion could be to stop at the first set of two thousand observations in which the number of increases and decreases are within 10\% of each other. He suggests that the maximum and minimum realized values could be used as bounds on the true value. Fox (1986) suggests that we compare the estimate of the integral based on a sample of \(2n\) points with the estimate based on \(n\) points and stop if the answer lies within some tolerance level. Paskov and Traub (1995) use a similar termination criterion based on successive errors: stop when the difference between two consecutive approximations using \(10,000i, i = 1, 2, \ldots, 1000\), sample points falls below some threshold. Owen (1994, 1995) proposes a hybrid of Monte Carlo and low-discrepancy methods which provides error estimates and has good convergence properties. In addition to these approaches, one can also run standard Monte Carlo at the outset and use the probabilistic error term to assess when enough low-discrepancy points have been used in the quasi-random calculation. This benchmarking with standard Monte Carlo would be useful if the same set of calculations were being carried out frequently with only slightly different input values. This situation is common in finance applications. There is often a need to perform the same set of calculations frequently, e.g., the risk analysis of a book of business at the end of each day. In these cases one can conduct experiments to see which sets of low-discrepancy sequences provide the
best results. The right number of low-discrepancy points could be determined just once at the outset.

Before leaving this section, we should mention some recent advances and new techniques to improve the performance of quasi-random Monte Carlo, Niederreiter and Xing (1995) and Ninomiya and Tezuka (1994) have proposed new low-discrepancy sequences that appear to have the potential to perform substantially better than previous methods. We have noted that the efficiency of quasi-random Monte Carlo improves as the integrand becomes smoother. Moskowitz and Caflisch (1995) illustrate procedures that can be used for this purpose. It is sometimes possible to enhance the performance of quasi-random sequences by reducing the effective dimension of the problem. Moskowitz and Caflisch also indicate how this can be accomplished in the discretization of a Wiener process and in the solution of the Feynman-Kac equation. This is relevant for finance applications since the prices of derivatives securities have a Feynman-Kac representation. See Aiworth, Broadie and Glasserman (1997), Berman (1996), and Caflisch, Morokoff and Owen for recent work applying low-discrepancy sequences with alternative constructions of Wiener processes. Spanier and Maize (1994) discuss a battery of techniques that can be used to improve the performance of quasi-Monte Carlo methods for relatively small sample sizes.

Next we compare the Monte Carlo method using pseudo-random numbers with the Faure, Halton, and Sobol' low-discrepancy methods.

3.1. Numerical results

For an initial comparison, we test the methods on the problem of pricing a European option on a single underlying asset with the usual Black-Scholes assumptions. In this framework, the Black-Scholes formula can be evaluated to give the true option values in order to compare alternative methods. Rather than using a single option, we evaluate the methods on a random sample of 500 options. The probability distribution of the parameters is chosen to represent a reasonable range of values in practical applications.\(^{20}\) The error measure that we use is root-mean-squared (RMS) relative error defined by

\[
RMS = \sqrt{\frac{1}{m} \sum_{i=1}^{m} \left( \frac{\hat{C}_i - C_i}{C_i} \right)^2},
\]

where \(i\) is the index of the \(m = 500\) options in the test set, \(C_i\) is the true option value, and \(\hat{C}_i\) is the estimated option value. The results are given in Fig. 1.

Fig. 1 plots RMS relative error against the number of points, \(n\). The Monte Carlo method (i.e., using pseudo-random numbers) displays the expected

\(^{20}\) The details of the distribution are given in Broadie and Detemple (1996).
Fig. 1. RMS relative error vs. number of points.

$O(1/\sqrt{n})$ convergence: e.g., increasing $n$ by a factor of 100 decreases the RMS error by a factor of 10. The low-discrepancy method using Faure sequences dominates the Monte Carlo method. Indeed, 129 Faure points gives an error lower than 1000 Monte Carlo points. The Sobol' method is the best of the three methods tested. Using 192 Sobol' points gives an error lower than 10,000 Monte Carlo points.

A major consideration in the comparison of methods is the overall computation time, not just the number of points. The Sobol' sequence numbers can be generated significantly faster than Faure numbers (see, e.g., Bratley and Fox 1988) and faster than most pseudo-random number methods. Hence, in the important RMS error versus computation time comparison, the relative advantage of the Sobol' method increases.

A low-discrepancy sequence will often have additional uniformity properties at certain points in the sequence (see, e.g., Fox 1986 and Bratley and Fox 1988). For example, in the Sobol' sequence the running average returns to 0.5 at the points $n = 2^k - 1$ for $k = 1, 2, \ldots$. One might expect that choosing $n$ to be one of these 'favorable' points would lead to better option price estimates. For large values of $n$, the advantage of using favorable points becomes negligible, but for small $n$ the effect can be quite significant. Indeed, in the experiment above, using
the Sobol' points 1–254 gives an RMS error of 10%, while using the points 1–255 gives an RMS error of 4%.\textsuperscript{21} Better results are often obtained by ignoring an initial portion of a low-discrepancy sequence. For example, using the Sobol' points 1–63 gives an RMS error of 13%, while using the Sobol' points 64–127 gives an RMS error of 2%. In the results in Fig. 1, the Sobol' sequence was always started at point 64, so the label 192 in Fig. 1 corresponds to the 192 Sobol' points from 64 to 255. Similarly, the Faure sequence was always started at point 16, so the label 129 in Fig. 1 corresponds to the 129 Sobol' points from 16 to 144.

3.2. One-dimensional vs. higher-dimensional sequences

It is sometimes asserted that low-discrepancy methods can be implemented in existing simulation programs by simply replacing the pseudo-random number generator with a low-discrepancy sequence generator. This naive approach can lead to disastrous results as the following example shows.

Consider pricing a European option on the maximum of two nondividend paying assets with the parameters: \( S_1 = S_2 = K = 100 \), \( \sigma_1 = \sigma_2 = 0.2 \), \( \rho = 0.3 \), \( r = 0.05 \), and \( T = 1 \). Under the usual Black-Scholes assumptions, a formula for the price of the option can be derived (see, e.g., Johnson 1987 or Stulz 1982) and gives a price of 16.442. Running one Monte Carlo simulation with 1000 points (hence 2000 random numbers) gave an estimated price of 16.279 with a standard error of 0.533. Using 2000 one-dimensional low-discrepancy values gave a price estimate of 4.320 using the Sobol' sequence and an estimate of 1.909 using the Faure sequence (starting at point 16). The cause of the problem can be seen by examining Figs. 2–5.

Figs. 2 and 3 show 1000 two-dimensional Faure and Sobol' points, respectively. The figures illustrate how the sequences fill the two-dimensional space in regular but different ways. By contrast, Figs. 4 and 5 show 2000 one-dimensional Faure and Sobol' points, respectively, plotted in two dimensions. The plots are created by taking successive points in the 1-dimensional sequence to be the \((x, y)\) coordinates in two-dimensional space. In neither figure are the points filling the two-dimensional space (note that the axes do not extend from 0 to 1) and this explains why the price estimates do not converge to the correct values. Even in the quarter of the unit square where the points fall, the points do not uniformly fill the space. This problem is reminiscent of the well-known 'collinearity' or 'hyperplane' problem of some pseudo-random number generators, but is even more serious with these low-discrepancy sequences.

A similar problem can occur if a high-dimensional low-discrepancy sequence is used for a problem of low dimension. Fig. 6 shows the 49th and 50th dimension of 1000 50-dimensional Faure points. Using the last two dimensions

\textsuperscript{21} We take the first point of the Sobol' sequence to be 0.5, not 0.0.
of the 50-dimensional sequence to price a two-dimensional option will give very poor results.

3.3. Higher-dimensional test

To test the effect of problem dimension, we price options in dimensions $d = 10, 50,$ and $100$. We price discretely sampled geometric average Asian options, because the problem dimension is easily varied and a closed form solution for the price is available (see Turnbull and Wakeman, 1991). The price
of a geometric average Asian option is given by

\[ C = E\left[e^{-rT}(\bar{S} - K)^+\right], \]

where \( \bar{S} = (\prod_{i=1}^{d} S_i)^{1/d} \) and \( S_i \) is the asset price at time \( iT/d \).

We test standard Monte Carlo, Monte Carlo with antithetic variates, and the low-discrepancy sequences of Faure, Sobol', and Halton.\(^{22}\) For each dimension,

\(^{22}\) We thank Spassimir Paskov and Joseph Traub for providing their code for the Sobol's sequences.
Fig. 6. Coordinates 49 and 50 of 1000 50-dimensional Faure points.

Fig. 7. Results with 50,000 points.

we select 500 option parameters at random, and compute RMS relative error (see Eq. (12)) for each method. Results for 50,000 and 200,000 sample points are given in Figs. 7 and 8, respectively. (The antithetic method uses 25,000 and 100,000 independent pairs of points, respectively).

\[^{23}\text{The details of the distribution are given in Broadie and Detemple (1996).}\]
Fig. 8. Results with 200,000 points.

Results for the Halton sequence were not competitive and are suppressed. RMS error for standard Monte Carlo is nearly independent of the problem dimension. The antithetic method gives minimal variance reduction. The relative advantage, in terms of RMS error, of the low-discrepancy sequences decreases with the problem dimension. For this test problem, the crossover point is beyond dimension 100.

4. Estimating price sensitivities

Most of the discussion in this paper centers on the use of Monte Carlo for pricing securities. In practice, the evaluation of price sensitivities is often as important as the evaluation of the prices themselves. Indeed, whereas prices for some securities can be observed in the market, their sensitivities to parameter changes typically cannot and must therefore be computed. Since price sensitivities are important measures of risk, the growing emphasis on risk management systems suggests a greater need for their efficient computation.

The derivatives of a derivative security's price with respect to various model parameters are collectively referred to as Greeks, because several of these are commonly referred to with the names of Greek letters.\(^\text{24}\) Perhaps the most important of these – and the one to which we give primary attention – is delta: the derivative of the price of a contingent claim with respect to the current price.

\(^{24}\)See, e.g., Chapter 14 of Hull (1997) for background.
of an underlying asset. The delta of a stock option, for example, is the derivative of the option price with respect to the current stock price. An option involving multiple underlying assets has multiple deltas, one for each underlying asset.

In the rest of this section, we discuss various approaches to estimating price sensitivities, especially delta. We begin by examining finite-difference approximations and show that these can be improved through the use of common random numbers. We then discuss direct methods that estimate derivatives without requiring resimulation at perturbed parameter values.

4.1. Finite-difference approximations

Consider the problem of computing the delta of the Black-Scholes price of a European call; i.e., computing

$$
\Delta = \frac{dC}{dS_0},
$$

where $C$ is the option price and $S_0$ is the current stock price. There is, of course, an explicit expression for delta, so simulation is not required, but the example is useful for purposes of illustration. A crude estimate of delta is obtained by generating a terminal stock price

$$
S_T = S_0e^{(r-(1/2)\sigma^2)T + \sigma \sqrt{T}Z}
$$

(see (2) for notation) from the current stock price $S_0$ and a second, independent terminal stock price

$$
S_T(\varepsilon) = (S_0 + \varepsilon)e^{(r-(1/2)\sigma^2)T + \sigma \sqrt{T}Z'}
$$

from the perturbed initial price $S_0 + \varepsilon$, with $Z$ and $Z'$ independent. For each terminal price, a discounted payoff can be computed like this:

$$
\tilde{C}(S_0) = e^{-rT} \max \{0, S_T - K\}, \quad \tilde{C}(S_0 + \varepsilon) = e^{-rT} \max \{0, S_T(\varepsilon) - K\}
$$

(see (3) for notation). A crude estimate of delta is then provided by the finite-difference approximation

$$
\tilde{\Delta} = \varepsilon^{-1} [\tilde{C}(S_0 + \varepsilon) - \tilde{C}(S_0)].
$$

By generating $n$ independent replications of $S_T$ and $S_T(\varepsilon)$ we can calculate the sample mean of $n$ independent copies of $\tilde{\Delta}$. As $n \to \infty$, this sample mean
converges to the true finite-difference ratio

\[ \varepsilon^{-1} [C(S_0 + \varepsilon) - C(S_0)], \]

where \( C(\cdot) \) is the option price as a function of the current stock price.

This discussion suggests that to get an accurate estimate of \( \Delta \) we should make \( \varepsilon \) small. However, because we generated \( S_T \) and \( S_T(e) \) independently of each other, we have

\[ \text{Var}[\tilde{\Delta}] = \varepsilon^{-2} (\text{Var}[\hat{C}(S_0 + \varepsilon)] + \text{Var}[C(S_0)]) = O(\varepsilon^{-2}), \]

so the variance of \( \tilde{\Delta} \) becomes very large if we make \( \varepsilon \) small. To get an estimator that converges to \( \Delta \) we must let \( \varepsilon \) decrease slowly as \( n \) increases, resulting in slow overall convergence. A general result of Glynn (1989) shows that the best possible convergence rate using this approach is typically \( n^{-1/4} \). Replacing the forward difference estimator in (15) with the central difference \( (2\varepsilon)^{-1} [\hat{C}(S_0 + \varepsilon) - \hat{C}(S_0 - \varepsilon)] \) typically improves the optimal convergence rate to \( n^{-1/3} \). These rates should be compared with \( n^{-1/2} \), the rate ordinarily expected from Monte Carlo.

Better estimators can generally be improved using the method of common random numbers, which, in this context, simply uses the same \( Z \) in (13) and (14). Denote by \( \tilde{\Delta} \) the finite-difference approximation thus obtained. For fixed \( \varepsilon \), the sample mean of independent replications of \( \tilde{\Delta} \) also converges to (16). The variance parameter is given by

\[ \text{Var}[\tilde{\Delta}] = \varepsilon^{-2} (\text{Var}[\hat{C}(S_0)] + \text{Var}[\hat{C}(S_0 + \varepsilon)] - 2\text{Cov}[\hat{C}(S_0), \hat{C}(S_0 + \varepsilon)]), \]

because \( \hat{C}(S_0) \) and \( \hat{C}(S_0 + \varepsilon) \) are no longer independent. Indeed, if they are positively correlated, then \( \tilde{\Delta} \) has smaller variance than \( \bar{\Delta} \). That they are in fact positively correlated follows from the monotonicity of the function mapping \( Z \) to \( \hat{C} \) by the argument used in our discussion of antithetics in Section 3. Thus, the use of common random numbers reduces the variance of the estimate of \( \Delta \).

The impact of this variance reduction is most dramatic when \( \varepsilon \) is small. A simple calculation shows that, using common random numbers,

\[ |\hat{C}(S_0 + \varepsilon) - \hat{C}(S_0)| \leq |S_T(e) - S_T| \leq \varepsilon e^{e^{-((1/2)\sigma^2)}T} + e^{(1/2)\sigma^2}T. \]

Because this upper bound has finite second moment, we may conclude that

\[ E[|\hat{C}(S_0 + \varepsilon) - \hat{C}(S_0)|^2] = O(\varepsilon^2), \quad (17) \]
and therefore that

$$\text{Var}[\varepsilon^{-1}(\hat{C}(S_0 + \varepsilon) - \hat{C}(S_0))] = O(1);$$

i.e., the variance of $\hat{A}$ remains bounded as $\varepsilon \to 0$, whereas we saw previously that the variance of $\hat{A}$ increases at rate $\varepsilon^{-2}$. Thus, the more precisely we try to estimate $A$ (by making $\varepsilon$ small) the greater the benefit of common random numbers. Moreover, this indicates that to get an estimator that converges to $A$ we may let $\varepsilon$ decrease faster as $n$ increases than was possible with $\hat{A}$, resulting in faster overall convergence. An application of Proposition 2 of L’Ecuyer and Perron (1994) shows that a convergence rate of $n^{-1/2}$ can be achieved in this case, and that is the best that can ordinarily be expected from Monte Carlo. For more on convergence rates using common random numbers see Glasserman and Yao (1992), Glynn (1989), and L’Ecuyer and Perron (1994).

The dramatic success of common random numbers in this example relies on the fast rate of mean-square convergence of $\hat{C}(S_0 + \varepsilon)$ to $\hat{C}(S_0)$ evidenced by (17). This rate does not apply in all cases. It fails to hold, for example, in the case of a digital option\(^{25}\) paying a fixed amount $B$ if $S_T > K$ and 0 otherwise. The price of this option is $C = e^{-rT}BP(S_T > K)$; the obvious simulation estimator is

$$\hat{C}(S_0) = I_{S_T > K}e^{-rT}B.$$ 

Because $\hat{C}(S_0)$ and $\hat{C}(S_0 + \varepsilon)$ differ only when $S_T \leq K < S_T(\varepsilon)$, we have

$$\text{E}[|\hat{C}(S_0 + \varepsilon) - \hat{C}(S_0)|^2] = B^2e^{-2rT}\text{P}(S_T \leq K < S_T(\varepsilon))$$

$$= B^2e^{-2rT}\text{P}(S_T \leq K < (1 + \varepsilon/S_0)S_T) = O(\varepsilon),$$

compared with $O(\varepsilon^2)$ for a standard call. As a result, delta estimation is more difficult for the digital option, and a similar argument applies to barrier options generally. Even in these cases, the use of common random numbers can result in substantial improvement compared with differences based on independent runs.

Table 6 compares the performance of four types of delta estimates: forward and central finite-differences with and without common random numbers. The methods are compared at four values of the perturbation parameter $\varepsilon$, and applied to the two options discussed above. The values in the table are estimated root mean square errors. The numerical results substantiate the analysis above. Much lower errors are obtained for the standard call than for the digital option, allowing for smaller $\varepsilon$; central differences beat forward differences; common random numbers help, but it helps the standard call more than the digital

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\(^{25}\) Also called a ‘binary’ or ‘cash-or-nothing’ option; see Hull (1997, p. 463).
Table 6
RMS errors for various delta estimation methods

<table>
<thead>
<tr>
<th>ε</th>
<th>Independent</th>
<th></th>
<th>Common</th>
<th></th>
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</thead>
<tbody>
<tr>
<td></td>
<td>Forward</td>
<td>Central</td>
<td>Forward</td>
<td>Central</td>
</tr>
<tr>
<td>Standard call option</td>
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<td>0.01</td>
<td>0.100</td>
</tr>
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<td></td>
<td>1</td>
<td>0.18</td>
<td>0.09</td>
<td>0.012</td>
</tr>
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<td></td>
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<td></td>
<td>0.01</td>
<td>7.47</td>
<td>8.98</td>
<td>0.006</td>
</tr>
<tr>
<td>Digital option</td>
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<td>0.51</td>
<td>0.37</td>
<td>0.51</td>
</tr>
<tr>
<td></td>
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<tr>
<td></td>
<td>5</td>
<td>0.16</td>
<td>0.07</td>
<td>0.11</td>
</tr>
<tr>
<td></td>
<td>1</td>
<td>0.67</td>
<td>0.34</td>
<td>0.14</td>
</tr>
</tbody>
</table>

Root mean square error of delta estimates for two options using four methods with various values of ε. Both options have $S_0 = 100$, $K = 100$, $\sigma = 0.40$, $r = 0.10$, and $T = 0.2$. The digital option has $B = 100$. Each entry is computed from 1,000 delta estimates, each estimate based on 10,000 replications. The value of delta is 0.580 for the first option and 2.185 for the second.

In several cases, the minimal error is obtained using a fairly large ε. This reflects the fact that the bias resulting from a large ε is sometimes overwhelmed by the large variance resulting from a small ε.

Although we have discussed common random numbers in only a limited context, it can easily be applied to a wide range of problems. If all stochastic inputs to a simulation are samples from the normal distribution, then common random numbers can be implemented by using the same samples at two different parameter settings. More generally, if the stochastic inputs are all drawn from a sequence of uniform random variates, then common random numbers can be implemented by using these variates at two different parameter settings.

4.2. Direct estimates

Even with the improvements in performance obtained from common random numbers, derivative estimates based on finite differences still suffer from two shortcomings. They are biased (since they compute difference ratios rather than derivatives) and they require multiple resimulations: estimating sensitivities to $d$ parameter changes requires repeatedly running one simulation with all parameters at their base values and $d$ additional simulations with each of the parameters perturbed. The computation of 10–50 Greeks26 for a single security

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26 Sensitivities to various changes in the yield curve often account for several of these.
is not unheard of, and this represents a significant computational burden when multiple resimulations are required.

Over the last decade, a variety of direct methods have been developed for estimating derivatives by simulation. Direct methods compute a derivative estimate from a single simulation, and thus do not require resimulation at a perturbed parameter value. Under appropriate conditions, they result in unbiased estimates of the derivatives themselves, rather than of a finite-difference ratio. Our discussion focuses on the use of pathwise derivatives as direct estimates, based on a technique generally called infinitesimal perturbation analysis (see, e.g., Glasserman, 1991).

The pathwise estimate of the true delta \( dC/dS_0 \) is the derivative of the sample price \( \hat{C} \) with respect to \( S_0 \). More precisely, it is

\[
\frac{d\hat{C}}{dS_0} = \lim_{\varepsilon \to 0} \left[ \hat{C}(S_0 + \varepsilon) - \hat{C}(S_0) \right],
\]

provided the limit exists with probability 1. If \( \hat{C}(S_o) \) and \( \hat{C}(S_o + \varepsilon) \) are computed from the same \( Z \), then provided \( S_T \neq K \), we have

\[
\frac{d\hat{C}}{dS_0} = \frac{d\hat{C}}{dS_T} \frac{dS_T}{dS_0} = e^{-rT} I_{[S_T > K]} \frac{S_T}{S_0}. \tag{18}
\]

We have used (13) to get

\[
\frac{dS_T}{dS_0} = e^{(r-(1/2)\sigma^2)T + \sigma \sqrt{T}Z} \frac{S_T}{S_0},
\]

and

\[
\frac{d\hat{C}}{dS_T} = e^{-rT} \frac{d}{dS_T} \max\{0, S_T - K\} = \begin{cases} e^{-rT}, & S_T > K, \\ 0, & S_T < K. \end{cases}
\]

At \( S_T = K \), \( C \) fails to be differentiable; however, since this occurs with probability zero, the random variable \( d\hat{C}/dS_0 \) is almost surely well defined.

The pathwise derivative \( d\hat{C}/dS_0 \) can be thought of as a limiting case of the common random numbers finite-difference estimator in which we evaluate the limit analytically rather than numerically. It is a direct estimator of the option delta because it can be computed directly from a simulation starting with \( S_0 \) without the need for a separate simulation at a perturbed value \( S_0 \). This is evident from the expression in (18). The question remains whether this estimator is unbiased; that is, whether

\[
E \left[ \frac{d\hat{C}}{dS_0} \right] = \frac{dC}{dS_0} = \frac{d}{dS_0} E[\hat{C}].
\]
The unbiasedness of the pathwise estimate thus reduces to the interchangeability of derivative and expectation. The interchange is easily justified in this case; see Broadie and Glasserman (1996) for this example and conditions for more general cases. Applying the same reasoning used above, we obtain the following pathwise estimators of three other Greeks for the Black-Scholes price:

$$\text{Rho}(dC/dr): \ K T e^{-rT} 1_{S_T \geq K};$$

$$\text{Vega}(dC/d\sigma): \ e^{-rT} 1_{S_T \geq K} \frac{S_T}{\sigma} (\ln(S_T/S_0) - (r - \frac{1}{2} \sigma^2) T),$$

$$\text{Theta}(-dC/dT): \ r e^{-rT} \max\{S_T - K, 0\} - 1_{S_T \geq K} e^{-rT} \frac{S_T}{2T} (\ln(S_T/S_0)$$

$$+ (r - \frac{1}{2} \sigma^2) T).$$

Each of these estimators is unbiased.

Of course, Monte Carlo estimators are not required for these derivatives because closed-form expressions are available for each. The Black-Scholes setting is useful for illustration, but the utility of the technique rests on its applicability to more general models. In Broadie and Glasserman (1996), pathwise estimates are derived and studied (both theoretically and numerically) for Asian options and a model with stochastic volatility. For example, the Asian-option delta estimate is simply

$$e^{-rT} \frac{\bar{S}}{S_0} 1_{\bar{S} > K},$$

where $\bar{S}$ is the average asset price used to determine the option payoff. Evaluating this expression takes negligible time compared with resimulating to estimate the option price from a perturbed initial stock price. The pathwise estimate is thus both more accurate and faster to compute than the finite-difference approximation. These advantages extend to a wide class of problems.

As already noted, the unbiasedness of pathwise derivative estimates depends on an interchange of derivative and expectation. In practice, this generally means that the security payoff should be a pathwise continuous function of the parameter in question. The standard call option payoff $e^{-rT} \max\{0, S_T - K\}$ is continuous in each of its parameters. An example where continuity fails is a digital option with payoff $e^{-rT} 1_{S_T > K} B$, with $B$ the amount received if the stock finishes in the money. 27 Because of the discontinuity at $S_T = K$, the

27 We used this example at the end of Section 3. The settings are related: problems for which common random numbers is particularly effective are generally problems to which the pathwise method can be applied even more effectively.
pathwise method (in its simplest form) cannot be applied to this type of option.

The problem of discontinuities often arises in the estimation of \textit{gamma}, the second derivative of an option price with respect to the current price of an underlying asset. Consider, again, the standard European call option. We have an expression for $d\hat{C}/dS_0$ in (18) involving the indicator $I_{\{S_T > K\}}$. This shows that $d\hat{C}/dS_0$ is discontinuous in $S_T$, preventing us from differentiating pathwise a second time to get a direct estimator of gamma.

To address the problem of discontinuities, Broadie and Glasserman (1996) construct \textit{smoothed} estimators. These estimators are unbiased, but not as simple to derive and implement as ordinary pathwise estimators. Broadie and Glasserman also investigate another technique for direct derivative estimation called the \textit{likelihood ratio method}. This method differentiates the probability density of an asset price, rather than the outcome of the asset price itself.\textsuperscript{28} The domains of this method and the pathwise method overlap, but neither contains the other. When both apply, the pathwise method generally has lower variance.

Overviews of these methods can be found in Glasserman (1991), Glynn (1987), and Rubinstein and Shapiro (1993). For discussions specific to financial applications see Broadie and Glasserman (1996) and Fu and Hu (1995).

5. Pricing American options by simulation

\textit{European} contingent claims have cash flows that cannot be influenced by decisions of the owner. Examples include European options, barrier options, and many types of swaps. By contrast, the cash flows of \textit{American} contingent claims depend both on the price path of the underlying asset or assets and the decisions of the owner. Many types of American contingent claims trade on exchanges and in the over-the-counter market. Examples include American options, American swaptions, shout options, and American Asian options. They also arise in other contexts, for example as \textquote{real options} in the theory of economic investment described in Dixit and Pindyck (1994).

To be concrete, suppose that we wish to estimate the quantity $\max_{\tau} E[e^{-r\tau} h(S_\tau)]$, where $r$ is the constant riskless interest rate, $h(S_\tau)$ is the payoff at time $\tau$ in state $S_\tau$, and the max is taken over all stopping times $\tau \leq T$. This formulation of the American pricing problem will suffice to illustrate the major points. First, note that the state can be vector-valued and hence applies to pricing American options on multiple assets. Second, since simulation algorithms are discrete in nature, the continuous-time exercise decision must be

\textsuperscript{28} Though not presented in a Monte Carlo context, the expressions in Carr (1993) are potentially relevant to this approach.
approximated by restricting the exercise opportunities to lie in a finite set of
times $0 = t_0 < t_1 < \cdots < t_d = T$. This is not always a serious restriction. For
example, for a call option on a stock which pays dividends at discrete points in
time, it can be shown that early exercise is only optimal just prior to the
ex-dividend dates. In other cases, Richardson or other extrapolation techniques
can be used to better approximate the price with exercise in continuous time
from a finite set of exercise opportunities. \footnote{Geske and Johnson (1984)
gave the first financial application of Richardson extrapolation. An
extensive treatment of extrapolation techniques is given in Marchuk and Shaidurov (1983).}
However, we now restrict attention to estimating the quantity
\begin{equation}
P = \max_{\tau} \mathbb{E}[e^{-r\tau}h(S_\tau)]. \tag{19}
\end{equation}
where the max is taken over all stopping times $\tau$ in the set $t_i$, for $i = 0, \ldots, d$. The
need to estimate an optimal stopping time is the crucial distinction between
American and European pricing problems.

If the state space is of low dimension, say three or less, a discretization scheme
together with a dynamic programming algorithm can often be used to numerically
approximate the value in (19). Even in these cases, simulation can be used to
estimate the expectation in the recursive step. Simulation-based methods
become essential when the dimension of the state space is large.

An obvious simulation-based algorithm for estimating the quantity in Eq. (19)
is to generate a random path of states $S_{i\tau}$, for $i = 1, \ldots, d$, and form the path
estimate
\begin{equation}
\hat{P} = \max_{i=0, \ldots, d} e^{-r\tau}h(S_i). \nonumber
\end{equation}
However, this estimator corresponds to using perfect foresight, and so it is biased
high. That is, $\mathbb{E}[\hat{P}] \geq P$, which follows immediately from the inequality
$\max_{i=0, \ldots, d} e^{-r\tau}h(S_i) \geq e^{-r\tau}h(S_i)$. A natural goal would be to develop an
alternative unbiased estimator. A negative result in this regard is provided in Broadie
and Glasserman (1997): among a large class of estimators, there is no unbiased
estimator of $P$. In particular, the estimators proposed in Tilley (1993), Grant et
al. (1994), and Barraquand and Martineau (1995) are all biased. Unfortunately,
they provide no way to estimate the extent of the bias or to correct for the bias in
a general setting. Broadie and Glasserman (1997) circumvent this problem by
developing two estimators, one biased high and one biased low (but both
asymptotically unbiased), which can be used together to form a valid confidence
interval for the quantity $P$. In the remainder of this section, we give brief
descriptions of the four methods mentioned and describe some strengths and weaknesses of each.

4.1. Tilley’s bundling algorithm

Tilley (1993) sparked considerable interest by demonstrating the potential practicality of applying simulation to pricing American contingent claims. Tilley describes a ‘bundling procedure’ for pricing an American option on a single underlying asset. To estimate \( P \) he suggests simulating \( n \) paths of asset prices denoted \( S_i(j) \) for \( i = 1, \ldots, d \) and \( j = 1, \ldots, n \) in the usual way. Next partition the asset price space and call the paths which fall into a given partition at a fixed time a ‘bundle’. A dynamic programming algorithm is applied to bundles to estimate \( C \). In particular, the estimated option price \( P_t(j) \) at time \( t \) for path \( j \) is the maximum of the immediate exercise value, \( h(S_t(j)) \), and the present value of continuing. The latter value is defined to be the average of \( e^{-r(t_{i+1} - t)} P_{t_{i+1}}(k) \) over all paths \( k \) which fall in the bundle containing path \( j \) at time \( t \). Details of the partitioning are given in Tilley (1993).

In order to implement the algorithm, all paths must be stored so they can be sorted into bundles at each time step. Since simulation typically requires a large number of paths for good estimates, the storage and sorting requirements can be significant. More importantly, the algorithm does not easily generalize to multiple state variables. In higher dimensions, it is not clear how to define the bundles. Even then it is likely that most partitions will contain very few paths and lead to a large bias, or the partitions will be so large that the continuation values are poorly estimated.

Because Tilley’s algorithm uses the same paths to estimate the optimal decisions and the value, the estimator tends to be biased high (although the bundling induces an approximation which is difficult to analyze). Tilley introduces a ‘sharp boundary’ variant which reduces the bias, but this variant does not easily generalize to higher dimensions. Carriere (1996) contains further analysis of Tilley’s algorithm and suggests a procedure based on spline functions to reduce the bias. It remains to be seen whether the spline procedure is practical for higher-dimensional problems. Nevertheless, for single state variable problems, Tilley demonstrated the potential practicality of applying simulation to American-type pricing problems.

4.2. Barraquand and Martineau’s stratified state aggregation (SSA) algorithm

Barraquand and Martineau (1995) propose a partitioning algorithm, but unlike Tilley’s bundling algorithm, they partition the payoff space instead of the state space. Hence, only a one-dimensional space is partitioned at each time
step, independent of the number of state variables. Their algorithm works as follows.

First, partition the payoff space into $K$ disjoint cells. Then simulate $n$ paths of asset prices denoted $S_i(j)$ for $i = 1, \ldots, d$ and $j = 1, \ldots, n$ in the usual way. For each payoff cell $k$ at time $t_i$, record the number of paths, $a_i(k)$, which fall into the cell. For each pair of cells $k$ and $l$ at consecutive times $t_i$ and $t_{i+1}$, record the number of paths, $b_i(k, l)$, which fall into both cells. Also, for each cell $k$ at time $t_i$, record the sum of the payoff values, $c_i(k) = \sum h(S_i(j))$, where the sum is over all paths $j$ which fall into cell $k$ at time $t_i$. The transition probability from $(t_i, k)$ to $(t_{i+1}, l)$ is approximated by $p_i(k, l) = b_i(k, l)/a_i(k)$. The estimated option price $P_i(k)$ at time $t_i$ in cell $k$ is the maximum of the immediate exercise value and the present value of continuing. The immediate exercise value is approximated by $c_i(k)/a_i(k)$. The present value of continuing is approximated by $e^{-r(t_{i+1}-t)} \sum_{l=1}^{K} p_i(k, l) P_{t_{i+1}}(l)$. This procedure can be applied backwards in time to determine the simulation estimate of the price $P$.

Details of a payoff space partitioning scheme are given in Barraquand and Martineau (1995). Once a single path is generated and the summary information $a, b$, and $c$ is recorded, the path can be discarded. Hence the storage requirements with this method are modest: on the order of $K^2 d$. One drawback of this method is a possible lack of convergence, as the following example illustrates.

Fig. 9 shows the evolution of two asset prices $(S_1, S_2)$. The option payoff is $h(S_1, S_2) = \max(S_1, S_2)$ and for convenience the riskless rate is taken to be zero. Using the risk-neutral probabilities in Fig. 9, the true value of the option at time $t_0$ is 11, which at time $t_1$ involves exercise in state (8, 4) but continuing in state (8, 8). (Note that asset 1 pays a dividend of 4 and asset 2 a dividend of 2 after time $t_1$.) When the states are partitioned by their payoffs, these two states are indistinguishable. As seen in the payoff evolution in Fig. 10, the best strategy at time $t_1$ in payoff state 8 is to continue. The apparent value of the option in Fig. 10 is 9 ($= (1/2)14 + (1/2)4$). In this example, partitioning the payoff space leads to a significant underestimate of the option value. Hence, a simulation algorithm based on partitioning the payoff space cannot converge to the correct value. Although this example may seem contrived, Broadie and Detemple (1994) show that the payoff value is not a sufficient statistic for determining the optimal exercise decision for options on the maximum of several assets. Indeed, the payoff process $h(S_t)$ is hardly ever Markovian.

There is currently no way to bound the error in the Barraquand and Martineau method. Without an error estimate, it is difficult to determine the

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30 In fact, they distinguish between partitioning the state space, which they term 'stratified state aggregation', and partitioning the payoff space, which they term 'stratified state aggregation along the payoff'. The latter method is the only one that they test or specify in detail. Hence we focus our discussion on this variant of their method.
appropriate number of paths to simulate or the appropriate number of partitions to use. Their method can be slightly modified to generate an option price estimate which is biased low as follows. Their procedure gives an exercise strategy based on the immediate exercise payoff. Using this strategy, a new (independent) set of paths can be simulated, and an option value can be estimated under the exercise strategy previously estimated. The resulting option price estimate will be biased low because the exercise policy is not, in general, the optimal policy. With this modification, the average direction of the error is known. Raymar and Zwecher (1996) extend the Barraquand and Martineau approach by basing the exercise decision on a partition of two state variables, rather than one.

4.3. Broadie and Glasserman’s algorithm

Broadie and Glasserman (1997) propose an algorithm based on simulated trees. In order to handle the bias problem, they develop two estimators, one
biased high and one biased low, but both convergent and asymptotically unbiased as the computational effort increases. A valid confidence interval for the true value $P$ is obtained by taking the upper confidence limit from the 'high' estimator and the lower confidence limit from the 'low' estimator. Briefly, their algorithm works as follows.

First, simulate a tree of asset prices (or, more generally, state variables) using $b$ branches at each node. Two paths emanating from a node evolve as independent copies of the state process. The high estimator, $\Theta$, is defined to be the value obtained by the usual dynamic programming algorithm applied to the simulated tree. Then repeat the process for $n$ trees, and compute a point estimate and confidence interval for $E[\Theta]$. A low estimator is obtained by modifying the dynamic programming algorithm at each node. Instead of using all $b$ branches to determine the decision and value, $b_1$ branches are used to determine the exercise decision, and the remaining $b_2 = b - b_1$ branches are used to determine the continuation value. Their actual low estimator, $\hat{\theta}$, includes another modification of this procedure which reduces the variance of the estimate. As before, estimates from $n$ trees are combined to give a point estimate and confidence interval for $E[\hat{\theta}]$. Details of the procedure can be found in Broadie and Glasserman (1997).

For the $\Theta$ estimator, all of the branches at a given node are used to determine the optimal decision and the corresponding node value, and this leads to an upward bias, i.e., $E[\Theta] \geq P$. For the $\theta$ estimator, the decision and the continuation value are determined from independent information sets. This eliminates the upward bias, but a downward bias occurs, i.e., $E[\hat{\theta}] \leq P$. The intuition for this result follows. If the correct decision is inferred at a node, the node value estimate would be unbiased. If the incorrect decision is inferred at a node, the node value estimate would be biased low because of the suboptimality of the decision. The expected node value is a weighted average of an unbiased estimate (based on the correct decision) and an estimate which is biased low (based on the incorrect decision). The net effect is an estimate which is biased low. Both estimators are consistent and asymptotically unbiased as $b$ increases.

The computational effort with this algorithm is order $nb^d$ and its main drawback is that $d$ cannot be too large for practical computations. Broadie and Glasserman (1997) give numerical results for options with $d = 4$. As mentioned earlier, to approximate option values with continuous exercise opportunities, some type of extrapolation procedure is required. Special care is necessary to implement extrapolation procedures within a simulation context because of the randomness in the estimates.

4.4. Other developments

Grant et al. (1994) describe a method specially designed to price American arithmetic Asian options on a single underlying asset. In this application the
optimal exercise decision depends on the current asset price and the current value of the average. Using repeated simulation runs, they attempt to identify the form of an optimal exercise policy based on these two pieces of information. Once an exercise policy is specified, simulation is used to estimate the option value under this fixed policy. Since the fixed policy is a suboptimal approximation to the optimal stopping rule, their procedure leads to a simulation estimator which is biased low.

GVW perform extensive sensitivity analysis which indicates that their option value estimate is relatively insensitive to deviations in the chosen exercise policy. So, it may be that their method gives good option price estimates relative to some accuracy level, but it is not clear how to quantify their error. It is not clear how to improve their estimates to an arbitrary accuracy level as the simulation effort increases. Their procedure is specific to the case of American Asian options and does not at this point constitute a general approach to pricing American contingent claims.

Bossaerts (1989) proposes two estimators of optimal early exercise, a moment estimator and a smooth optimization estimator, and studies their convergence properties. His method appears to require a parametric representation of the exercise boundary and may therefore face difficulties in higher dimension. The optimization approach described in Fu and Hu (1995) also requires a parametric representation.

Rust (1995) studies the general problem of solving discrete decision problems, which include optimal stopping problems as a special case. He develops a Monte Carlo method and shows that it succeeds in breaking the ‘curse of dimensionality’ in these problems. Rust’s focus is on computational complexity, but his approach appears to provide a promising direction for finance applications.

4.5. Summary

The valuation of securities with American-type features requires the determination of optimal decisions. High dimension versions of these problems arise from multiple state variables and/or path dependencies. Although simulation is a powerful tool for solving some higher-dimensional problems, conventional wisdom was that simulation could not be applied to American-style pricing problems. The algorithms described here represent the first attempts to solve these problems that were long thought to be computationally intractable.

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31 We thank A. Dixit for pointing us to this reference.
6. Further topics

We conclude this paper with a brief mention of two important areas of current work in the application of Monte Carlo methods to finance, not discussed in this article.

A central numerical issue in simulating interest rates, asset prices with stochastic volatilities, and other complex diffusions is the accurate approximation of stochastic differential equations by discrete-time processes. Kloeden and Platen (1992) discuss a variety of methods for constructing discrete-time approximations with different orders of convergence. Andersen (1995) applies some of these to interest-rate models. In general, decreasing the time increment in a discrete approximation can be expected to give more accurate results, but at the expense of greater computational effort. Duffie and Glynn (1995) analyze this trade-off and characterize asymptotically optimal time steps as the overall computational effort grows.

In this article we have focused almost exclusively on the use of Monte Carlo for pricing. A related, growing area of application is risk management – in particular, the use of Monte Carlo to assess value at risk, credit risk, and related measures. For some examples of recent applications in these areas see Iben and Brotherton-Ratcliffe (1994), Lawrence (1994), and Beckström and Campbell (1995).

Appendix: Moment controls beat moment matching asymptotically

As mentioned in Section 2.4, any time a moment is available for use with moment matching, it can alternatively be used as a control variate. In this appendix, we argue that moment matching is asymptotically equivalent to a control variate technique with suboptimal coefficients, and is therefore dominated by the optimal use of moments as controls. This asymptotic link applies in large samples. A related link between linear and nonlinear control variates is made in Glynn and Whitt (1989), but the current setting does not fit their framework.

Let $Z_1, Z_2, \ldots$ be i.i.d. (not necessarily normal) with mean $\mu$ and variance $\sigma^2$. Let $s$ denote the sample standard deviation of $Z_1, \ldots, Z_n$ and $\bar{Z}$ their sample mean. Suppose we want to estimate $E[f(Z)]$ for some function $f$. The standard estimator is $n^{-1} \sum_{i=1}^k f(Z_i)$ and the moment matching estimator is $n^{-1} \sum_{i=1}^k f(\bar{Z}_i)$ with $\bar{Z}_i$ defined in $(9)$. For each $i$, the scaled difference

$$\sqrt{n}(\bar{Z}_i - Z_i) = \sqrt{n} \left( \frac{\sigma - s}{s} \right) Z_i - \sqrt{n} [\sigma \bar{Z}_i / s - \mu]$$
converges in distribution, by the central limit theorem for \( \hat{Z} \) and \( s \). Thus, \( (\hat{Z} - Z_i) = O_p(n^{-1/2}) \) (see, e.g., Appendix A of Pollard 1984 for \( O_p, o_p \) notation).

Suppose now that, with probability one, \( f \) is differentiable at \( Z_i \). Then

\[
 f(\hat{Z}_i) = f(Z_i) + f'(Z_i)[\hat{Z}_i - Z_i] + o_p(n^{-1/2}),
\]

suggesting that up to terms \( o_p(n^{-1/2}) \) the moment matching estimator and standard estimator are related via

\[
 \frac{1}{n} \sum_{i=1}^{n} f(\hat{Z}_i) \approx \frac{1}{n} \sum_{i=1}^{n} f(Z_i) + \frac{1}{n} \sum_{i=1}^{n} f'(Z_i)[\hat{Z}_i - Z_i]
\]

\[
 = \frac{1}{n} \sum_{i=1}^{n} f(Z_i) + \frac{1}{n} \sum_{i=1}^{n} f'(Z_i) \left( \frac{\sigma}{s} - 1 \right) Z_i - \frac{\sigma}{s} \hat{Z} + \mu \left( \frac{\sigma}{s} - 1 \right)
\]

\[
 + \left( \frac{1}{n} \sum_{i=1}^{n} f'(Z_i) \right) \left( \beta \sigma - \frac{\sigma}{s} \hat{Z} \right)
\]

\[
 = \frac{1}{n} \sum_{i=1}^{n} f(Z_i) + \beta_1 \left( \frac{\sigma}{s} - 1 \right) + \beta_2 \left( \mu - \frac{\sigma}{s} \hat{Z} \right)
\]

where \( \hat{\beta}_i \to \beta_i, i = 1, 2, \) as \( n \to \infty \), with

\[
 \beta_1 = \mathbb{E}[f'(Z)Z] \quad \text{and} \quad \beta_2 = \mathbb{E}[f'(Z)].
\]

Thus, moment matching is asymptotically equivalent to using

\[
 \left( \frac{\sigma}{s} - 1 \right) \quad \text{and} \quad \left( \mu - \frac{\sigma}{s} \hat{Z} \right)
\]

as controls (both quantities converge to zero almost surely) with estimates of coefficients \( \beta_1, \beta_2 \). In general, these do not coincide with the optimal coefficients \( \beta_1^*, \beta_2^* \), so moment matching is asymptotically dominated by the control variate method. In addition, the controls in (20) introduce some bias (as does moment matching itself) because though they converge to zero they do not have mean zero for finite \( n \). In contrast, the more natural moment control variates \((s^2 - \sigma^2)\) and \((\hat{Z} - \mu)\) have mean zero for all \( n \) and thus introduce no bias.
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