A Comparison of Iterated Optimal Stopping and Local Policy Iteration for American Options Under Regime Switching

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Received: 6 September 2012 / Revised: 9 May 2013 / Accepted: 22 May 2013 © Springer Science+Business Media New York 2013

Abstract A theoretical analysis tool, iterated optimal stopping, has been used as the basis of a numerical algorithm for American options under regime switching (Le and Wang in SIAM J Control Optim 48(8):5193–5213, 2010). Similar methods have also been proposed for American options under jump diffusion (Bayraktar and Xing in Math Methods Oper Res 70:505–525, 2009) and Asian options under jump diffusion (Bayraktar and Xing in Math Fin 21(1):117–143, 2011). An alternative method, local policy iteration, has been suggested in Huang et al. (SIAM J Sci Comput 33(5):2144–2168, 2011), and Salmi and Toivanen (Appl Numer Math 61:821–831, 2011). Worst case upper bounds on the convergence rates of these two methods suggest that local policy iteration should be preferred over iterated optimal stopping (Huang et al. in SIAM J Sci Comput 33(5):2144–2168, 2011). In this article, numerical tests are presented which indicate that the observed performance of these two methods is consistent with the worst case upper bounds. In addition, while these two methods seem quite different, we show that either one can be converted into the other by a simple rearrangement of two loops.

Keywords Iterated optimal stopping · Local policy iteration · Regime switching

Mathematics Subject Classification 60G40 · 65N06 · 65K15

This work was supported by Tata Consulting Services and the Natural Sciences and Engineering Research Council of Canada.

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Published online: 02 June 2013



1 Introduction

In this work, our focus is on comparing iterative methods for solving discretized Hamilton Jacobi Bellman equations. As an example, we consider the case of an American option, assuming that the underlying asset follows a regime switching model. This gives rise to a coupled system of variational inequalities (VIs). After discretizing these equations, using a fully implicit or Crank–Nicolson finite difference scheme, the main challenge is efficiently solving the resulting nonlinear algebraic equations.

We center our comparison on two methods, *iterated optimal stopping* and *local policy iteration*. Iterated optimal stopping was developed in a recent article [27], where the authors analyse the properties of the solution of a finite time optimal stopping (American) option pricing problem under regime switching. That particular work is based on similar analysis carried out in [4] for American options under jump diffusion, and in [5] for Asian options under jump diffusion. The method of analysis in [27] constructs a sequence of functions which converge monotonically to the exact solution of the optimal stopping problem. In [27], a numerical method is developed based in this concept. The iterated optimal stopping technique was also used to develop numerical algorithms for American and Asian options under jump diffusion [4,5].

In [29], a local policy iteration method was developed for solving the discretized variational inequality for pricing American options under a jump diffusion process. This method was generalized for the case of American options under regime switching in [21]. In addition, a form of local policy iteration was suggested in [31], however no analysis of the convergence of this iteration was given.

Superficially, iterated optimal stopping and local policy iteration appear to be very different algorithms. However, when viewed in more general terms, these two techniques are simply different iterative methods for solving the same set of discretized nonlinear algebraic equations. In [21], worst case upper bounds on the convergence rate for both methods were developed. The bound for local policy iteration is favorable compared to the bound for iterated optimal stopping. However, it is not clear that the upper bounds in [21] are tight. In addition, no numerical tests using iterated optimal stopping are given in [21].

In this paper we provide an extensive set of numerical tests to study the performance of iterated optimal stopping and local policy iteration. Our tests show that the performance of both algorithms is consistent in practice with the worst case upper bounds. However it is also the case that these methods appear to approach their worst case bounds only for long term contracts. We also observe that implementation is not an issue. Indeed, we show that an existing implementation of iterated optimal stopping can be converted to local policy iteration by effectively interchanging two loops. Finally, iterated optimal stopping requires significantly more storage than local policy iteration. As a result we recommend the use of local policy iteration.

The remainder of the paper proceeds as follows. Section 2 gives the basics of the regime switching model, including the nonlinear partial differential equation for pricing American options. The continuous version of the iterated optimal stopping method is presented in Sect. 3 while the discretization of the continuous American option problem is presented in Sect. 4. Section 5 provides the matrix representation of the nonlinear discrete equations with iterative methods for solving these equations given in the following section, including algorithm formulations of local policy iteration and iterated optimal stopping. Numerical tests appear in Sect. 7 followed by a concluding section.



2 Regime Switching Model

Financial practitioners are intimately aware of the shortcomings of the now ubiquitous Black-Scholes framework. Asset prices driven by a geometric Brownian motion process with constant volatility and drift cannot replicate the distribution of returns seen in historical stock prices [30]. A regime switching process is a computationally parsimonious technique for introducing stochastic volatility into the underlying stochastic model. Regime switching processes have been applied to problems in electricity markets [6], long term insurance guarantees [19], forestry valuation [13], and gas storage [14].

Define a finite set of K regimes and a volatility σ_j and drift rate $\mu_j^{\mathbb{P}}$ associated with each state $j = 1 \dots K$. The \mathbb{P} denotes that drift rates are observed in the real-world measure. A continuous Markov chain process is used to transition between any two states. The system of stochastic equations governing the regime switching process is

$$dS = \mu_j^{\mathbb{P}} S dt + \sigma^j S dZ + \sum_{k=1}^K (\xi_{jk} - 1) S dX_{jk}; \ j = 1, \dots, K,$$
 (2.1)

where S is the asset price, dZ is the increment of a Wiener process and ξ_{jk} is the jump amplitude when a transition from state $j \to k$ occurs. The Poisson process dX_{jk} is defined by

$$dX_{jk} = \begin{cases} 1 \text{ with probability} & \lambda_{jk}^{\mathbb{P}} dt + \delta_{jk} \\ 0 \text{ with probability} & 1 - \lambda_{jk}^{\mathbb{P}} dt - \delta_{jk} \end{cases}$$

$$\lambda_{jk}^{\mathbb{P}} \geq 0 \; ; \; j \neq k$$

$$\lambda_{jj}^{\mathbb{P}} = -\sum_{\substack{k=1\\k \neq j}}^{K} \lambda_{jk}^{\mathbb{P}}. \tag{2.2}$$

where λ_{jk} is the probability of transitioning from regime $j \to k$. Observe that the asset price jumps from $S \to \xi_{jk} S$ when such a transition occurs. To prevent a jump in the absence of a regime switch, set $\xi_{jj} = 1$.

To determine the fair-value of an option under regime switching, the hedging portfolio in regime j, P_j is constructed

$$P_{j} = -\mathcal{V}_{j} + eS + \sum_{k=1}^{K-1} w_{k} F_{k}$$
 (2.3)

where V_j is the no-arbitrage value of the option in regime j, e is the amount held of the underlying asset with price S, and w_k is the amount held of the additional hedging instruments with price F_k . Provided that the underlying asset and the additional hedging instruments form a non-redundant set, all risk can be eliminated through a dynamic hedging strategy [26]. This permits the use of a no-arbitrage argument leading to the American option pricing equations

$$\min \left[\frac{\partial \mathcal{V}_j}{\partial \tau} - \mathcal{L}_j \mathcal{V}_j - \lambda_j \mathcal{J}_j \mathcal{V}, \quad \mathcal{V}_j - \mathcal{V}^* \right] = 0; \quad j = 1, \dots, K,$$
 (2.4)

where $\tau = T - t$ represents time running backwards from expiry T, and $\mathcal{V}^*(S) = \mathcal{V}(S, \tau = 0)$ is the payoff condition. The differential operators \mathcal{L}_i and \mathcal{J}_i are defined as

$$\mathcal{L}_{j}\mathcal{V}_{j} = \left(\frac{\sigma_{j}^{2}S^{2}}{2}\right)\frac{\partial^{2}\mathcal{V}_{j}}{\partial S^{2}} + (r - \rho_{j})S\frac{\partial\mathcal{V}_{j}}{\partial S} - (r + \lambda_{j})\mathcal{V}_{j}$$
 (2.5a)

$$\mathcal{J}_{j}\mathcal{V} = \sum_{\substack{k=1\\k\neq j}}^{K} \frac{\lambda_{jk}}{\lambda_{j}} \mathcal{V}_{k}(\xi_{jk}S, \tau) , \qquad (2.5b)$$

where r > 0 is the risk-free rate (i.e. the rate of return of a security which has no possibility of default) and

$$\rho_{j} = \sum_{\substack{k=1\\k\neq j}}^{K} \lambda_{jk} (\xi_{jk} - 1) \; ; \; \lambda_{j} = \sum_{\substack{k=1\\k\neq j}}^{K} \lambda_{jk}.$$
 (2.6)

It is assumed that ξ_{jk} are deterministic functions of S and t. The risk neutral transition densities λ_{jk} are not unique. In practice, we calibrate the parameters in Eq. (2.5) to market data, consistent with the market's pricing measure. Note that Eq. (2.5a) is independent of $\mu_j^{\mathbb{P}}$, due to the usual hedging arguments [26].

For computational purposes, Eq. (2.4) will be posed on the localized domain

$$(S, \tau) \in [0, S_{\text{max}}] \times [0, T].$$
 (2.7)

No boundary condition is required at S = 0 while at $S = S_{\text{max}}$, a Dirichlet condition is imposed (in this paper we use the payoff)

$$\mathcal{V}(S_{\text{max}}, \tau) = \mathcal{V}^*(S_{\text{max}}), \quad \forall \tau. \tag{2.8}$$

We truncate any jumps which would require data outside the computational domain. The error in this approximation is small in regions of interest if S_{max} is sufficiently large [26]. More precisely, the term $\mathcal{V}_k(\xi_{jk}S,\tau)$ in Eq. (2.5b) is replaced by $\mathcal{V}_k(\min(S_{\text{max}},\xi_{jk}S),\tau)$.

Remark 2.1 (Viscosity Solution) Equation (2.4) is a special case of the more general systems of VIs considered in [16], where it is shown that VIs such as (2.4) have unique, continuous, locally bounded viscosity solutions in the interior of the solution domain. The definition of a viscosity solution must be generalized for systems of PDEs [9,16,23,24]. One of the advantages of the viscosity solution approach is that boundary and initial conditions can be interpreted in the relaxed viscosity sense, which means that the solution may not converge to the initial condition in the usual sense [2,25]. For example, [15] discusses the case of discontinuous initial conditions which are common in financial applications (i.e. digital options).

Remark 2.2 (Localized Boundary Conditions) The effect of the localized boundary condition (2.8) can be made arbitrarily small by selecting S_{max} sufficiently large [3].

3 Iterated Optimal Stopping: Continuous Case

The basic idea of iterated optimal stopping [4,5,27] is to consider a sequence of functions $(\mathcal{V}_i)^m$, such that

$$\min \left[\frac{\partial (\mathcal{V}_j)^{m+1}}{\partial \tau} - \mathcal{L}_j(\mathcal{V}_j)^{m+1} - \lambda_j \mathcal{J}_j(\mathcal{V})^m, (\mathcal{V}_j)^{m+1} - \mathcal{V}^* \right] = 0 \; ; \; j = 1, \dots, K.$$
(3.1)

Under suitable assumptions, it can be shown that the sequence $(V_j)^m$ converges monotonically to the solution of Eq. (2.4) [27].



Note that at each iteration m in Eq. (3.1), the problem reduces to solving a set of decoupled American type problems, one for each regime. This allows the authors in [27] to deduce properties of the solution V_j . The operator \mathcal{J}_j is replaced by an integral operator in [4,5] and similar arguments are constructed for jump diffusions.

Remark 3.1 (Interpretation of Eq. (3.1)) Problem (3.1) effectively reduces problem (2.4) to a sequence of simpler problems. Each of these simpler problems has known properties. This makes formulation (3.1) attractive from a mathematical point of view. However, we will see that formulation (3.1) is not well-suited to numerical computation.

4 Discretization

Numerically solving the American option pricing equations in (2.4) is achieved through a finite-difference discretization. A price grid is constructed consisting of a set of i_{max} nodes

$$\{S_1, S_2, \dots, S_{i_{\max}}\}\$$
 (4.1)

following a sequence of L timesteps of size $\Delta \tau$

$$\{\tau^0, \tau^1, \dots, \tau^L\} \tag{4.2}$$

where $\tau^n = n \Delta \tau$.

Denote the numerical solution at (S_i, τ^n) in regime j by $V_{i,j}^n$. A vector of solution values is constructed as follows

$$V^{n} = \left[V_{1,1}^{n}, \dots, V_{i_{\max},1}^{n}, \dots, V_{1,K}^{n}, \dots, V_{i_{\max},K}^{n} \right]'. \tag{4.3}$$

The solution vector has length $N = K \cdot i_{\text{max}}$. For succinctness, a single row index is often used to refer to entries of V^n as follows

$$V_{\ell}^{n} = V_{i,j}^{n}; \ \ell = (j-1)i_{\text{max}} + i.$$
 (4.4)

4.1 L Operator Discretization

Let \mathcal{L}_{j}^{h} denote the discrete form of \mathcal{L}_{j} in (2.5a). We use a combination of central, forward and backward differencing to generate a discretization of the form

$$(\mathcal{L}_{j}^{h}V^{n})_{ij} = \alpha_{i,j}V_{i-1,j}^{n} + \beta_{i,j}V_{i+1,j}^{n} - (\alpha_{i,j} + \beta_{i,j} + r + \lambda_{j})V_{i,j}^{n}$$
(4.5)

Central differencing is used as much as possible with forward backward differencing used only as required to ensure that the positive coefficient condition

$$\alpha_{i,j} \ge 0; \quad \beta_{i,j} \ge 0; \quad \forall i, j,$$
 (4.6)

is enforced. We refer the reader to [18] for details.

4.2 \mathcal{J} Operator Discretization

Let \mathcal{J}_j^h denote the discrete form of \mathcal{J}_j in (2.5b). This operator is discretized through a linear interpolation approximation as follows

$$[\mathcal{J}_j^h V^n]_{i,j} = \sum_{\substack{k=1\\k\neq j}}^K \frac{\lambda_{jk}}{\lambda_j} \mathcal{I}_{i,j,k}^h V^n, \tag{4.7}$$

where

$$\mathcal{I}_{i,j,k}^{h} V^{n} = w V_{m,k}^{n} + (1 - w) V_{m+1,k}^{n}$$

$$\simeq \mathcal{V}_{k}(\min(S_{i_{max}}, \xi_{jk} S_{i}), \tau^{n}), \tag{4.8}$$

with $w \in [0, 1]$ such that

$$\min(S_{i_{\max}}, \xi_{jk}S_i) = wS_m + (1 - w)S_{m+1}. \tag{4.9}$$

Note that jumps which extend outside the price grid are approximated by the value at the largest node $S_{i_{max}}$.

4.3 Imposing the American Constraint

The American constraint is handled implicitly by rewriting (2.4) in direct control form as

$$\max_{\varphi \in \{0,1\}} \left[\Omega \varphi (\mathcal{V}^* - \mathcal{V}_j) - (1 - \varphi) \left(\frac{\partial \mathcal{V}_j}{\partial \tau} - \mathcal{L}_j \mathcal{V}_j - \lambda_j \mathcal{J}_j \mathcal{V} \right) \right] = 0, \tag{4.10}$$

where we have introduced a scaling parameter $\Omega > 0$. In exact arithmetic, the introduction of the scaling factor Ω in Eq. (4.10) has no effect. However, any iterative algorithm used to solve Eq. (4.10) requires comparison to two terms which have different units. In order to remedy this situation, we use the scaling suggested in [21]

$$\Omega = \frac{C}{\Delta \tau},\tag{4.11}$$

where C > 0 is a dimensionless constant.

Equation (4.10) is discretized using fully-implicit ($\theta = 1$) or Crank–Nicolson ($\theta = 0.5$) timestepping

$$(1 - \varphi_{i,j}^{n+1}) \left(\frac{V_{i,j}^{n+1}}{\Delta \tau} - \theta \mathcal{L}_{j}^{h} V_{i,j}^{n+1} \right) + \Omega \varphi_{i,j}^{n+1} V_{i,j}^{n+1}$$

$$= \left(1 - \varphi_{i,j}^{n+1} \right) \frac{V_{i,j}^{n}}{\Delta \tau} + \Omega \varphi_{i,j}^{n+1} \mathcal{V}_{i}^{*} + \left(1 - \varphi_{i,j}^{n+1} \right) \lambda_{j} \theta \left[\mathcal{J}_{j}^{h} V^{n+1} \right]_{i,j}$$

$$+ \left(1 - \varphi_{i,j}^{n+1} \right) (1 - \theta) \left[\mathcal{L}_{j}^{h} V_{i,j}^{n} + \lambda_{j} \left[\mathcal{J}_{j}^{h} V^{n} \right]_{i,j} \right]; i < i_{\text{max}}$$

$$V_{i,j}^{n+1} = \mathcal{V}_{i}^{*}; i = i_{\text{max}}, \tag{4.12a}$$

where

$$\begin{aligned} \{\varphi_{i,j}^{n+1}\} &\in \operatorname*{arg\ max}_{\varphi \in \{0,1\}} \left\{ \Omega\ \varphi\left(\mathcal{V}_{i}^{*} - V_{i,j}^{n+1}\right) - (1-\varphi) \left(\frac{V_{i,j}^{n+1} - V_{i,j}^{n}}{\Delta \tau} - \theta\left(\mathcal{L}_{j}^{h} V_{i,j}^{n+1} + \lambda_{j} \left[\mathcal{J}_{j}^{h} V^{n+1}\right]_{i,j}\right) - (1-\theta) \left(\mathcal{L}_{j}^{h} V_{i,j}^{n} + \lambda_{j} \left[\mathcal{J}_{j}^{h} V^{n}\right]_{i,j}\right) \right) \right\}, \end{aligned} \tag{4.12b}$$

and $\varphi_{\ell} = 1$ indicates early exercise is optimal at node ℓ , otherwise $\varphi_{\ell} = 0$.

Remark 4.1 (Convergence to the viscosity solution) It is straightforward to show, using the methods in [18] that scheme (4.12), is unconditionally l_{∞} stable ($\theta = 1$), monotone and consistent, and hence converges to the viscosity solution of Eq. (2.4).



5 General Form of Equations

Let Q be a vector of controls defined by

$$Q = \left[\varphi_{1,1}, \dots, \varphi_{i_{\max},1}, \dots, \varphi_{1,K}, \dots, \varphi_{i_{\max},K}\right]'. \tag{5.1}$$

The discrete Eqs. (4.12) can then be written as a nonlinear matrix problem

$$\mathcal{A}^*(Q) U = \mathcal{C}(Q, V^n)$$
with $Q_{\ell} = \underset{Q_s \in \{0, 1\}}{\arg \max} \left[-\mathcal{A}^*(Q)U + \mathcal{C}(Q, V^n) \right]_{\ell}$ (5.2)

where U is the desired solution V^{n+1} at the next time level and A^* is an $N \times N$ matrix. For computational purposes, we split the A^* matrix as follows

$$A^*(O) = A(O) - B(O) \tag{5.3}$$

with $\mathcal{A}(Q)$ containing the coefficients of nodes coupled within the same regime, while $\mathcal{B}(Q)$ contains the coefficients that couple nodes in different regimes. This splitting will be useful when constructing numerical algorithms to solve the nonlinear equations in Sect. 6.

The matrix coefficients for nodes $i < i_{\text{max}}$ can be deduced from Eqs. (4.12) as shown below

$$[\mathcal{A}(Q)U]_{\ell} = [\mathcal{A}U]_{\ell} = (1 - \varphi_{\ell}) \left(\frac{U_{\ell}}{\Delta \tau} - \theta \mathcal{L}_{j}^{h} U_{\ell} \right) + \varphi_{\ell} \Omega U_{\ell}$$

$$[\mathcal{B}(Q)U]_{\ell} = [\mathcal{B}U]_{\ell} = (1 - \varphi_{\ell}) \lambda_{j} \theta \left[\mathcal{J}_{j}^{h} U \right]_{\ell}$$

$$[\mathcal{C}(Q, V^{n})]_{\ell} = C_{\ell} = (1 - \varphi_{\ell}) \frac{V_{\ell}^{n}}{\Delta \tau} + \varphi_{\ell} \Omega \mathcal{V}_{i}^{*}$$

$$+ (1 - \varphi_{\ell})(1 - \theta) \left[\mathcal{L}_{j}^{h} V_{\ell}^{n} + \lambda_{j} [\mathcal{J}_{j}^{h} V^{n}]_{\ell} \right]. \tag{5.4}$$

The boundary condition at node $i = i_{max}$ requires that

$$[\mathcal{A}U]_{\ell} = U_{i_{\max},j} \; ; \; [\mathcal{B}U]_{\ell} = 0 \; ; \; \mathcal{C}_{\ell} = \mathcal{V}^*_{i_{\max}} \; ; \; \ell = (j-1)i_{\max} + i_{\max}.$$
 (5.5)

6 Iterative Methods

Two numerical methods are presented to solve the nonlinear matrix problem in (5.2). A local policy iteration is compared to an iterated optimal stopping iteration proposed by [4]. It will be useful to note the following result.

Lemma 6.1 (Uniqueness of Solution of (5.2)) There exists a unique solution to Eq. (5.2).

Proof Note that the positive coefficient condition (4.6) guarantees that $\mathcal{A}^*(Q)$ is an M-matrix. In addition, the construction of $\mathcal{C}(Q, V^n)$ in Eq. (5.4) ensures that $\mathcal{C}(Q, V^n)$ is bounded for any finite mesh. Existence and uniqueness follow from the results in [7,22].

6.1 Local Policy Iteration

In this approach, we directly discretize Eq. (2.4). Following the approach of [31], the matrix splitting in (5.3) is used to solve the American control problem with the regime coupling terms lagged behind one iteration. This local policy method is outlined in Algorithm 6.1.



Algorithm 6.1 Local Policy Iteration

```
1: (V^0)^0 = \text{payoff}

2: for n = 0, 1, 2, ..., L - 1 do

3: (V^{n+1})^0 = V^n

4: for k = 0, 1, 2, ... until converge do

5: Solve: \max_{Q_{\ell} \in \{0, 1\}} \left\{ -\mathcal{A}(Q)(V^{n+1})^{k+1} + \mathcal{B}(Q)(V^{n+1})^k + \mathcal{C}(Q, V^n) \right\} = 0

6: if k > 0 and \max_{\ell} \frac{|(V_{\ell}^{n+1})^{k+1} - (V_{\ell}^{n+1})^k|}{\max\left[scale, |(V_{\ell}^{n+1})^{k+1}|\right]} < tolerance_{outer} then

7: V^{n+1} = (V^{n+1})^{k+1}; break from the iteration

8: end if

9: end for

10: end for
```

This method was suggested in [29] for American options under jump diffusion. Note that we write Algorithm 6.1 for all timesteps, although the actual iteration occurs only within each timestep. This will facilitate comparison with the iterated optimal stopping algorithm. We remark that a related idea for impulse control problems was suggested in [10–12].

The relative convergence tolerance in Line 5 in Algorithm 6.1 uses a *scale* factor to ensure that unrealistic accuracy criteria is not required for very small option values. Typically scale = 1 for options priced in dollars. Line 5 in Algorithm 6.1 is solved by iterating on $(V^{n+1})^{k+1}$. We use Algorithm 6.2 to solve for $(V^{n+1})^{k+1}$.

Algorithm 6.2 Inner Iteration (Policy Iteration)

```
1: U^{0} = (V^{n+1})^{k}
2: for m = 0, 1, 2, ... until converge do
3: Q^{m} = \underset{Q_{\ell} \in \{0,1\}}{\operatorname{arg max}} \left\{ -A(Q)U^{m} + B(Q)(V^{n+1})^{k} + C(Q, V^{n}) \right\}
4: Solve: A(Q^{m})U^{m+1} = B(Q^{m})(V^{n+1})^{k} + C(Q^{m}, V^{n})
5: if k > 0 and \underset{\ell}{\operatorname{max}} \frac{|U_{\ell}^{m+1} - U_{\ell}^{m}||}{\operatorname{max} \left[scale, |U_{\ell}^{m+1}||\right]} < tolerance_{inner} then
6: break from the iteration
7: end if
8: end for
9: (V^{n+1})^{k+1} = U^{m+1}
```

For American options, Line 5 in Algorithm 6.1 reduces to solving a linear complementarity problem (LCP). In general, even for a tridiagonal LCP problem, an iterative method is required [17] for the local American problem. In the special case of a simple put or call, only a single iteration is necessary [8], since the exercise region is simply connected to the boundary. Consequently, for a simple put or call, it would always be more efficient to use the direct Brennan and Schwartz method [8] to solve the local American problem, as in [29], for the local policy iteration. However, the standard Brennan and Schwartz algorithm [8] cannot be directly applied to more general problems, such as the American butterfly, which we will use as a test case.

Theorem 6.1 Suppose $(V^{n+1})^k$ is the exact solution of Line 5 in Algorithm 6.1, and let $E^k = (V^{n+1})^k - V^{n+1}$ where V^{n+1} is the exact solution to Eq. (5.2). If the matrices A and B are given by Eq. (5.4) then the local policy iteration in Algorithm 6.1 converges at the rate



$$\frac{\parallel E^{k+1} \parallel_{\infty}}{\parallel E^{k} \parallel_{\infty}} \le \frac{\theta \hat{\lambda} \Delta \tau}{1 + \theta (r + \hat{\lambda}) \Delta \tau} \quad \text{where } \hat{\lambda} = \max_{j} \lambda_{j}. \tag{6.1}$$

Remark 6.1 Result (6.1) is independent of the technique used to solve Line 5 in Algorithm 6.1.

Remark 6.2 In Theorem 6.1, we have assumed that $(V^{n+1})^k$ is the exact solution of Line 5 in Algorithm 6.1. In general, this will not be the case if an iterative method such as Algorithm 6.2 is used to solve Line 5 in Algorithm 6.1. A more detailed estimate for the convergence rate taking this into account is given in "Appendix".

6.2 Iterated Optimal Stopping: Discretization

The philosophy behind this approach is to discretize Eq. (3.1). In [27], the authors numerically solve a sequence of iterated optimal stopping problems to price an American put under a regime switching model. This *global-in-time* iteration is seen in Algorithm 6.3. Line 4 is solved in the same manner as described in Algorithm 6.2.

Algorithm 6.3 Iterated Optimal Stopping

```
1: (V^n)^0 = \text{payoff}; \ n = 0, ..., L

2: \mathbf{for} \ k = 0, 1, 2, ... \ \text{until converge do}

3: \mathbf{for} \ n = 0, 1, ..., L - 1 \ \mathbf{do}

4: \mathbf{Solve} : \max_{Q_\ell \in \{0,1\}} \left[ -A(Q)(V^{n+1})^{k+1} + B(Q)(V^{n+1})^k + C(Q, (V^n)^{k+1}) \right] = 0

5: \mathbf{end} \ \mathbf{for}

6: \mathbf{if} \ k > 0 \ \text{and} \ \max_{\ell, n} \frac{|(V_\ell^{n+1})^{k+1} - (V_\ell^{n+1})^k|}{\max \left[ scale, |(V_\ell^{n+1})^{k+1}| \right]} < tolerance_{outer} \ \mathbf{then}

7: \mathbf{break} \ \mathbf{from} \ \mathbf{the} \ \mathbf{iteration}

8: \mathbf{end} \ \mathbf{if}

9: \mathbf{end} \ \mathbf{for}
```

By storing the solutions at all timesteps in a single vector and constructing two $N(L+1) \times N(L+1)$ matrices, Algorithm 6.3 can be rewritten in the same form as Algorithm 6.1. As with Theorem 6.1 we can determine the following convergence bound for the iterated optimal stopping iteration.

Theorem 6.2 Assume that $(V^{n+1})^k$ is the exact solution to Line 4 in Algorithm 6.3. Let $E^k = \max_n \| (V^{n+1})^k - V^{n+1} \|_{\infty}$ where V^{n+1} is the exact solution to Eq. (5.2) at timestep n+1. If the matrices A and B are given by Eq. (5.4) and fully implicit time stepping $(\theta = 1)$ is used, then the iterated optimal stopping iteration converges at the rate

$$\frac{E^{k+1}}{E^k} \le \left(1 - \frac{1}{\left[1 + \Delta\tau(\hat{\lambda} + r)\right]^L}\right) \left(\frac{\hat{\lambda}}{\hat{\lambda} + r}\right) \tag{6.2}$$

Proof See [21]. \Box



Remark 6.3 As the timestep size decreases, the global convergence bound (6.2) becomes

$$\lim_{\substack{\Delta\tau\to 0\\L\to\infty\\L\Delta\tau=T}} \left(1 - \left[1 + \Delta\tau(\hat{\lambda} + r)\right]^{-L}\right) \left(\frac{\hat{\lambda}}{\hat{\lambda} + r}\right) = \left(1 - e^{-T(\hat{\lambda} + r)}\right) \left(\frac{\hat{\lambda}}{\hat{\lambda} + r}\right). \tag{6.3}$$

Hence, timestep refinements will only have a limited impact on the convergence of the iterated optimal stopping iteration. Contracts spanning longer periods of time have a theoretically worse convergence bound. Additionally, storage of solution vectors at every time level is required.

The iterated optimal stopping convergence bound in (6.2) is worse than the local policy bound in (6.1) since

$$\frac{\hat{\lambda}\Delta\tau}{1+(r+\hat{\lambda})\Delta\tau} \le \left(1-\left[1+\Delta\tau(\hat{\lambda}+r)\right]^{-L}\right)\left(\frac{\hat{\lambda}}{\hat{\lambda}+r}\right) . \tag{6.4}$$

Proposition 6.1 (Extension to Crank–Nicolson Case) Provided that

$$\Delta \tau < 2(\Delta \tau)_{\varrho} \tag{6.5}$$

numerical tests	
0 1 0	
On each refinement	a new grid

Table 1 Grid and timestep refinement levels used during

On each refinement, a new grid point is placed halfway between all old grid points and the number of timesteps is doubled. A constant timestep size is used

Refinement	Timesteps	America	an put	American butterfly	
		Nodes	Unknowns	Nodes	Unknowns
0	25	29	58	51	153
1	50	57	114	101	303
2	100	113	226	201	603
3	200	225	450	401	1203
4	400	449	898	801	2403
5	800	897	1794	1601	4803
6	1,600	1, 793	3, 586	3, 201	9, 603
7	3, 200	3, 585	7, 170	6, 401	19, 203

Table 2 American put: convergence as the grid and timesteps tend to zero

Refinement	$(\sigma_1, \sigma_2) = (0.3, 0.5)$)	$(\sigma_1, \sigma_2) = (0.5, 0.4)$			
	Value (regime 1)	Value (regime 2)	Value (regime 1)	Value (regime 2)		
0	0.596066588	1.220613143	1.247602448	0.933862220		
1	0.623233337	1.231744588	1.258122381	0.951342835		
2	0.627143687	1.236397726	1.263093502	0.955611738		
3	0.629053164	1.238462238	1.265163537	0.957249428		
4	0.629544512	1.239313670	1.266042273	0.957915213		
5	0.629756273	1.239696598	1.266438090	0.958209403		
6	0.629846395	1.239878850	1.266626785	0.958348152		
7	0.629888511	1.239968194	1.266719449	0.958416059		
Le and Wang	0.6300	1.2381	1.2647	0.9586		

Iterated optimal stopping is used with fully implicit timesteps. Parameter values are r=0.2, T=1, Strike=10, $\lambda_{1,2}=0.05$, $\lambda_{2,1}=0.15$. Jump amplitudes are 1 and option values are at S=10. Le and Wang results are from [27]



Table 3 Input parameters and data for the American put problem

Strike (put), K	10
Risk free rate, r	0.05
Maximum grid price, $S_{i_{max}}$	500
Transition probability $1 \rightarrow 2, \lambda_{1,2}$	3
Transition probability $2 \rightarrow 1, \lambda_{2,1}$	2
Jump amplitudes, $\xi_{1,2}$, $\xi_{2,1}$	1.0, 1.0
Volatilities, σ_1 , σ_2	0.3, 0.4

Table 4 American put—refinement results for the local policy iteration using fully implicit timesteps

Refinement	Value	Outer iterations per timestep	Inner iterations per outer iteration	Normalized CPU time
T=1				
0	1.149561815	6.00	2.10	1
1	1.165728338	5.00	2.12	2
2	1.171152560	4.00	2.11	4
3	1.173223677	3.21	2.11	8
4	1.174102036	3.00	2.11	22
5	1.174505970	3.00	2.09	68
6	1.174699595	2.45	2.11	200
7	1.174795968	2.05	2.11	713
T = 10				
0	2.480493130	15.6	2.06	3
1	2.527224461	10.4	2.07	4
2	2.545702990	7.28	2.10	7
3	2.552125529	5.45	2.12	13
4	2.554365849	4.38	2.15	30
5	2.555244216	3.76	2.16	84
6	2.555626744	3.07	2.18	248
7	2.555804716	3.00	2.18	925

Option values are at S=10 in Regime 1. 1 unit of CPU time is based on level zero, T=1. Compare with Table 5

where $(\Delta \tau)_e$ is the maximum stable explicit timestep size, then we can extend the above convergence analysis for the Crank Nicolson case $(\theta = 1/2)$

$$\frac{E^{k+1}}{E^k} \le \left(1 - \left[\frac{1 - (1-\theta)\Delta\tau(\hat{\lambda} + r)}{1 + \theta\Delta\tau(\hat{\lambda} + r)}\right]^L\right) \left(\frac{\hat{\lambda}}{\hat{\lambda} + r}\right). \tag{6.6}$$

Proof This follows the same steps as used in [21].

Remark 6.4 In the limit as $\Delta t \rightarrow 0$, then bound (6.6) becomes

$$\lim_{\substack{\Delta\tau\to 0\\L\to\infty\\L\to\tau=T}} \left(1 - \left[\frac{1 - (1-\theta)\Delta\tau(\hat{\lambda}+r)}{1 + \theta\Delta\tau(\hat{\lambda}+r)}\right]^L\right) \left(\frac{\hat{\lambda}}{\hat{\lambda}+r}\right) = \left(1 - e^{-T(\hat{\lambda}+r)}\right) \left(\frac{\hat{\lambda}}{\hat{\lambda}+r}\right). \tag{6.7}$$



Table 5 American put—refinement results for the iterated optimal stopping iteration using fully implicit timesteps

Refinement	Value	Outer iterations	Inner iterations per timestep per outer iteration	Normalized CPU time
T=1				
0	1.149561951	14	1.98	4
1	1.165728413	14	1.98	8
2	1.171152525	13	2.04	17
3	1.173224277	13	2.12	43
4	1.174102017	13	2.28	125
5	1.174505855	13	2.52	417
6	1.174700149	13	2.89	1, 583
7	1.174796138	13	3.46	6, 809
T = 10				
0	2.480497995	59	1.90	15
1	2.527228409	54	1.90	30
2	2.545706581	51	2.01	67
3	2.552129517	49	2.17	167
4	2.554369087	49	2.46	498
5	2.555247451	48	2.98	1, 739
6	2.555628049	48	3.86	7, 290
7	2.555804652	48	5.15	35, 274

Option values are at S = 10 in Regime 1. CPU times are normalized so that the time for local policy iteration, level zero is 1 unit (see Table 4). Compare with Table 4

This has the same limit as $\Delta \tau \to 0$ as in Eq. (6.3). Condition (6.5) is required to ensure that iterated optimal stopping form of the global matrix is an M matrix [21]. This is a very severe condition in practice, and is not practically useful (that is, it would be better to simply use an explicit method in this case).

Remark 6.5 (Rate of Convergence) A similar bound on the rate of convergence for the case of an Asian option under jump diffusion was obtained in [5], which is a special case of the more general result in [21]. It is straightforward to obtain obtain a similar convergence estimate for American Asian options, again using the method in [21]. In [5], this is referred to as *exponential convergence*. In the usual terminology of iterative methods, this would be referred to as linear convergence.

In [27], the rather weaker estimate

$$\frac{E^{k+1}}{E^k} \le \left(\frac{\hat{\lambda}}{\hat{\lambda} + r}\right) \tag{6.8}$$

was obtained.

Proposition 6.2 (Uniqueness of Solution: Algorithms 6.1 and 6.3) If the matrix $A^*(Q)$ is an M-matrix, then if local policy iteration and iterated optimal stopping converge, they converge to the same solution.



Table 6 American put—refinement results for the local policy iteration using Crank–Nicolson timesteps

Refinement	Value	Outer iterations	Inner iterations per timestep per outer iteration	Normalized CPU time
T=1				
0	1.158047827	5.08	2.09	1
1	1.170297442	4.06	2.13	2
2	1.173605701	3.69	2.11	4
3	1.174518713	3.02	2.09	8
4	1.174777835	3.00	2.08	23
5	1.174855570	3.00	2.06	73
6	1.174879679	2.15	2.04	191
7	1.174888084	2.01	2.07	713
T = 10				
0	2.494986976	11.4	2.07	2
1	2.535628033	7.80	2.08	3
2	2.550159207	5.94	2.11	6
3	2.554443725	4.79	2.12	12
4	2.555558848	4.02	2.15	30
5	2.555855693	3.18	2.16	80
6	2.555939143	3.01	2.15	266
7	2.555963088	2.98	2.15	1,015

Option values are at S=10 in Regime 1. 1 unit of CPU time is based on level zero, T=1. Compare with Table 7

Proof From Algorithm 6.1 and Algorithm 6.3, we have that at convergence

$$\mathcal{A}^*(Q) V^{n+1} = \mathcal{C}(Q, V^n)$$
with $Q_{\ell} = \underset{Q_s \in \{0,1\}}{\arg \max} \left[-\mathcal{A}^*(Q) V^{n+1} + \mathcal{C}(Q, V^n) \right]_{\ell}$
(6.9)

From Lemma 6.1, the solution of Eq. (6.9) is unique.

Remark 6.6 (Implementation Issues) The same algorithm is used to solve Line 5 in Algorithm 6.1 as used in Line 4 in Algorithm 6.3. Consequently, Algorithm 6.3 can be converted to Algorithm 6.1 by simply interchanging the n and k loops. This is a trivial implementation change, which has a significant effect on the properties of the algorithm.

Remark 6.7 (Jump Diffusions) The iterated optimal stopping algorithm has been proposed as a method for pricing American options under jump diffusions [4]. The general analysis in [21] can be used to obtain convergence rate estimates for local policy iteration and iterated optimal stopping for this problem as well. Similar to the regime switching case, the convergence bounds for iterated optimal stopping compare unfavorably with local policy iteration.

7 Numerical Examples

The comparably poor upper bound on the convergence rate and heavy memory requirements would suggest that iterated optimal stopping is less computationally efficient



 Table 7
 American put—refinement results for the iterated optimal stopping iteration using Crank–Nicolson timesteps

Refinement	Value	Outer iterations per timestep	Inner iterations per outer iteration	Normalized CPU time
T=1				
0	1.158047915	11	2.04	3
1	1.170297512	10	2.04	6
2	1.173605959	10	2.04	14
3	1.174518781	10	2.06	35
4	1.174777806	10	2.15	100
5	1.174855530	10	2.34	331
6	1.174879899	10	2.59	1, 233
7	1.174888119	10	3.01	5, 286
T = 10				
0	2.494988694	37	2.02	10
1	2.535630069	33	1.99	20
2	2.550160719	32	2.01	44
3	2.554444966	31	2.16	111
4	2.555558751	30	2.42	325
5	2.555856723	30	2.83	1, 139
6	2.555939003	30	3.44	4, 597
7	2.555962940	30	4.40	21, 570

Option values are at S = 10 in Regime 1. CPU times are normalized so that the time for local policy iteration, level zero is 1 unit (see Table 6). Compare with Table 6

compared to local policy iteration. Nonetheless, several authors have used this type of iteration recently [4,5,27]. We therefore provide numerical comparisons of the local policy and iterated optimal stopping iterations in this section to verify that the observed performance of these two algorithms is consistent with the worst case upper bounds. Two numerical examples are treated in this section, an American put contract based on the data used in [27], and an American butterfly contract based on data from [21].

The sequence of grid and timestep refinements can be seen in Table 1. The price grids consist of a fine mesh nearby the strikes and an increasingly coarser grid further away from the strike. On each grid refinement, new fine grid nodes are placed between each two coarse grid nodes, and the timesteps are doubled. A direct method is used to solve the linear tridiagonal matrix in Line 4 of Algorithm 6.2.

A scale factor of $\Omega = 10^6/\Delta \tau$ and outer convergence tolerance of 10^{-6} is used for all experiments (Line 6 in Algorithms 6.1 and 6.3). The inner iteration tolerance in Algorithm 6.2 was 10^{-9} .

7.1 Validation

In this section we show the convergence of the discretization as the grid is refined. Table 2 shows results for two of the American put cases reported in [27]. This is a two regime example. We use fully implicit timestepping with iterated optimal stopping. The parameter values for these results are shown in the table caption. Our results are comparable with those in [27]. However, there is no report of grid/timestep size sensitivity or convergence criteria in [27].



Table 8 American butterfly—refinement results for the local policy iteration using fully implicit timesteps

Refinement	Value	Outer iterations per timestep	Inner iterations per outer iteration	Normalized CPU time
T = 0.5				
0	4.414610687	5.64	2.14	1
1	4.437635032	4.96	2.15	2
2	4.450644836	4.00	2.19	5
3	4.456010904	3.47	2.21	13
4	4.458284154	3.02	2.24	42
5	4.459342999	3.00	2.24	156
6	4.459853289	3.00	2.23	717
7	4.460104008	2.26	2.23	2710
T = 10				
0	8.503526680	20.0	2.04	3
1	8.391584342	12.6	2.07	5
2	8.371971829	8.60	2.10	9
3	8.370349958	6.35	2.12	22
4	8.370656431	5.10	2.15	64
5	8.371011766	4.12	2.19	199
6	8.371245326	3.42	3.42 2.23	
7	8.371375743	3.05	2.24	3414

Option values are at S=93 in Regime 2. CPU times are normalized so that the time for local policy iteration, level zero is 1 unit. Compare with Table 9

7.2 American Put

A two-state model with no jumps in the asset price is used to price an American put whose payoff is given by $V^* = \max(K - S, 0)$ where K is the strike. The model parameters are shown in Table 3.

Remark 7.1 (Termination of Iteration) Note that the termination criteria in Line 6 of Algorithm 6.1 and Algorithm 6.3 do not precisely correspond to the bounds in Eqs. (6.1) and (6.2), since in practice we do not have the exact solution available. The termination criteria in Algorithms 6.1 and 6.3 can be regarded as practical implementations. However, we expect (as will be verified in the numerical tests) that the trends in the number of iterations will be roughly consistent with the bounds in Eqs. (6.1) and (6.2).

Refinement results using the local policy iteration with fully implicit and Crank–Nicolson timesteps are shown in Tables 4, 6 respectively. The number of iterations required for convergence is reported for various grid/timestep refinements and contract lengths.

Similar results using the iterated optimal stopping iteration are in Tables 5, 7. The table heading "Inner Iterations" refers to those iterations required to solve Lines 5 and 4 in Algorithms 6.1 and 6.3 respectively.

Remark 7.2 (Condition (6.5)) For all the Crank–Nicolson tests, we violate condition (6.5). This has no effect on the local policy iteration algorithm, but convergence of



Table 9 American butterfly—refinement results for the iterated optimal stopping iteration using fully implicit timesteps

Refinement	Value	Outer iterations	Inner iterations per timestep per outer iteration	Normalized CPU time
T = 0.5				
0	4.414610844	12	2.21	3
1	4.437635066	12	2.37	7
2	4.450644870	12	2.65	21
3	4.456011081	12	3.10	73
4	4.458284196	12	3.76	288
5	4.459343003	12	4.77	1, 295
6	4.459853288	12	6.35	7, 618
7	4.460104147	12	8.74	45, 699
T = 10				
0	8.503527951	71	1.99	16
1	8.391586147	65	2.08	36
2	8.371973661	62	2.26	99
3	8.370351440	60	2.67	328
4	8.370657097	59	3.35	1, 290
5	8.371012428	58	4.56	6, 015
6	8.371246794	58	6.28	36, 450
7	8.371376073	58	8.76	22, 2080

Option values are at S = 93 in Regime 2. CPU times are normalized so that the time for local policy iteration, level 0 is 1 unit (see Table 8). Compare with Table 8

the iterated optimal stopping algorithm cannot be guaranteed if condition (6.5) is not satisfied.

As expected from the convergence bound in (6.1), the local policy method requires fewer outer iterations as the grid and timesteps are refined. This holds for both the shorter T=1 contract and the longer T=10 contract. Hence when solving larger problems, fewer outer iterations are needed.

In contrast, the iterated optimal stopping iteration exhibits several undesirable characteristics. The number of outer iterations approaches a fixed non-zero value as the grid and timestep is refined. This value gets increasingly larger for longer contract lengths in accordance with (6.3). The number of inner iterations increases with the refinement level. This is a bit surprising, since the same algorithm is used to solve Line 5 in Algorithm 6.1 and Line 4 in Algorithm 6.3. A detailed examination revealed a large increase in the number of inner iterations within the first few outer iterations as grid refinements were made. This may be due to the poor initial estimate for the solution in the case of iterated optimal stopping. As a result, the CPU run times grow quite rapidly with every grid refinement as compared to the local policy run times.

Remark 7.3 (Complexity of Both Methods) Since Algorithm 6.2 is the same for both methods, the ratio of the number of floating point operations for both methods is



Table 10	American	butterfly-	-refinement	results	for	the	local	policy	iteration	using	Crank-Nicolson
timesteps											

Refinement	Value	Outer iterations per timestep	Inner iterations per outer iteration	Normalized CPU time
T = 0.5				
0	4.440162686	5.08	2.12	1
1	4.451422139	4.06	2.15	2
2	4.457862480	3.94	2.15	6
3	4.459717103	3.08	2.17	15
4	4.460172252	3.01	2.17	52
5	4.460299471	3.00	2.18	197
6	4.460336267	2.61	2.06	767
7	4.460347552	2.09	2.21	3, 123
T = 10				
0	8.521803224	14.0	2.05	2
1	8.400544555	9.40	2.09	4
2	8.376389091	6.84	2.11	9
3	8.372576548	5.21	2.14	24
4	8.371770123	4.25	2.17	70
5	8.371587805	3.84	2.19	244
6	8.371527683	3.10	2.22	947
7	8.371517241	3.01	2.22	4, 226

Option values are at S = 93 in Regime 2. CPU times are normalized so that the time for local policy iteration, level zero is 1 unit. Compare with Table 11

Flops for Iterated Optimal Stopping

Flops for Local Policy Iteration

$$= \frac{\text{Outer Iterations} \times \text{Inner Iterations} : \text{Iterated Stopping}}{\text{Outer Iterations} \times \text{Inner Iterations} : \text{Local Policy}}.$$
 (7.1)

The ratio of CPU times in the Tables is approximately equal to the ratio of complexities in Eq. (7.1), for the finer grids (the timings are not accurate for the coarse grids).

7.3 American Butterfly

An American butterfly contract is priced with a payoff given by

$$V^* = \max(S - K_1, 0) - 2\max(S - (K_1 + K_2)/2, 0) + \max(S - K_2, 0)$$
 (7.2)

where K_1 and K_2 represent the two strikes. Data and parameter values for the butterfly example are shown below. We assume the existence of an American contract with payoff (7.2), which can only be early exercised as a unit. This contract has been used as severe test case by several authors [1,20,28].



Refinement	Value	Outer iterations	Inner iterations per timestep per outer iteration	Normalized CPU time
T = 0.5				
0	4.440162700	10	2.17	3
1	4.451422164	9	2.33	7
2	4.457862497	9	2.56	19
3	4.459717136	9	2.90	66
4	4.460172252	9	3.42	256
5	4.460299466	9	4.22	1, 144
6	4.460336333	9	5.39	6, 429
7	4.460347592	9	7.18	36, 853
T = 10				
0	8.521803835	44	2.13	12
1	8.400545174	40	2.19	28
2	8.376389721	38	2.41	78
3	8.372577132	37	2.73	258
4	8.371770660	36	3.33	999
5	8.371588095	36	4.11	4, 494
6	8.371527985	36	5.32	25, 490
7	8.371517265	36	7.20	14, 8238

Table 11 American butterfly—refinement results for the iterated optimal stopping iteration using Crank–Nicolson timesteps

Option values are at S = 93 in Regime 2. CPU times are normalized so that the time for local policy iteration, level zero is 1 unit (see Table 10). Compare with Table 10

$$K_{1} = 90 \; ; \quad K_{2} = 110 \quad ; \quad r = 0.02 \; ; \quad S_{i_{\text{max}}} = 5000$$

$$\lambda = \begin{bmatrix} -3.2 & 0.2 & 3.0 \\ 1.0 & -1.08 & .08 \\ 3.0 & 0.2 & -3.2 \end{bmatrix} \; ; \quad \xi = \begin{bmatrix} 1.0 & 0.90 & 1.1 \\ 1.2 & 1.0 & 1.3 \\ 0.95 & 0.8 & 1.0 \end{bmatrix} \; ; \quad \sigma = \begin{bmatrix} .2 \\ .15 \\ .30 \end{bmatrix} . \tag{7.3}$$

Note that in this case, the solutions will not have continuous $(V_j)_S$. The payoff is also not convex, hence this precludes the analysis in [27]. However, this does not cause any difficulty if we pose the solution to problem (2.4) in terms of viscosity solutions.

Refinement results for the butterfly payoff using the local policy iteration with fully implicit and Crank–Nicolson timestepping are displayed in Tables 8, 10 respectively. Accompanying iterated optimal stopping refinement results are in Tables 9, 11. Note that the same issues affecting the iterated optimal stopping method discussed in Sect. 7.2 persist when using the second order Crank–Nicolson discretization. The method requires significantly longer runtimes as compared to the local policy iteration, particularly for highly refined grids and longer contract lengths.

In view of Remark 7.1, we carried out additional tests to check the convergence bound in (6.2) for the American Butterfly example. In these tests, an approximation to the successive error ratio E^{k+1}/E^k was computed. The exact option values are approximated with a numerical solution generated using a convergence tolerance of 10^{-12} . The approximate error terms



Table 12 Approximate error
ratios, E^{k+1}/E^k , for the iterated
optimal stopping iteration

Table 12 Approximate error ratios, E^{k+1}/E^k , for the iterated optimal stopping iteration	Refinement	Convergence bounds		Approximate error ratio optimal stopping	
		Local policy	Optimal stopping	Avg value	Max value
	T = 0.5				
	0	0.060	0.785	0.241	0.349
	1	0.031	0.790	0.231	0.365
	2	0.016	0.793	0.224	0.362
	3	0.008	0.794	0.221	0.362
American Butterfly case. The	4	0.004	0.794	0.219	0.362
approximate error ratio values are computed using fully implicit	T = 10				
timesteps. The convergence	0	0.559	0.994	0.819	0.940
bounds also correspond to fully	1	0.389	0.994	0.793	0.925
implicit timesteps (see (6.1) and (6.2)). The exact solution	2	0.242	0.994	0.779	0.922
was approximated by generating	3	0.138	0.994	0.771	0.920
a solution with a convergence tolerance of 10^{-12}	4	0.074	0.994	0.767	0.920

 E^k were then calculated at each iteration and timestep. The average and maximum ratio of successive error estimates for a fully implicit iterated optimal stopping iteration are shown in Table 12. The observed error ratios tend to the bound (6.2) as the expiry time increases.

8 Conclusions

Analysis of the iterated optimal stopping algorithm proposed in [27] for American options under regime switching shows that this technique is inferior in terms of worst case bounds on the rate of convergence compared with local policy iteration [29,21]. Numerical tests indicate that the observed performance of both these methods is consistent with the bounds on the convergence rates, using a common type of convergence test. In addition, iterated optimal stopping requires considerably more storage compared to local policy iteration.

We also note that iterated optimal stopping has been proposed for jump diffusion problems [4,5]. The analysis of the convergence rates for these problems is similar to that for regime switching, and can be easily carried out using the general form for the discretized equations as discussed in [21]. The analysis shows that the convergence bounds for iterated optimal stopping are worse than for local policy iteration.

Since it is a simple matter to convert an iterated optimal stopping implementation to local policy iteration (this simply involves interchanging two loops), we strongly recommend use of local policy iteration.

Appendix: Error Bound for Local Policy Iteration with Inexact Inner Solution

In this Appendix, we generalize the result in Theorem 6.1 to include the effect of an approximate solution to Line 5 in Algorithm 6.1. We need only generalize the steps used in [21].



If V^{n+1} is a solution to Eq. (5.2) then

$$\max_{Q'} \left\{ -\mathcal{A}(Q')V^{n+1} + \mathcal{B}(Q')V^{n+1} + \mathcal{C}(Q', V^n) \right\} = 0, \tag{9.1}$$

while from Algorithm 6.1, we have

$$\max_{Q} \left\{ -\mathcal{A}(Q)(V^{n+1})^{k+1} + \mathcal{B}(Q)(V^{n+1})^{k} + \mathcal{C}(Q, V^{n}) \right\} = \mathcal{E}^{k}. \tag{9.2}$$

The term \mathcal{E}^k in Eq. (9.2) takes into account that that we may not necessarily have the exact solution to Line 5 in Algorithm 6.1, if we use Algorithm 6.2.

Subtracting Eq. (9.1) from Eq. (9.2) we obtain

$$\mathcal{E}^{k} = \max_{Q} \left\{ -\mathcal{A}(Q)(V^{n+1})^{k+1} + \mathcal{B}(Q)(V^{n+1})^{k} + \mathcal{C}(Q, V^{n}) \right\} - \max_{Q'} \left\{ -\mathcal{A}(Q')V^{n+1} + \mathcal{B}(Q')V^{n+1} + \mathcal{C}(Q', V^{n}) \right\} \leq \max_{Q} \left\{ -\mathcal{A}(Q)((V^{n+1})^{k+1} - V^{n+1}) + \mathcal{B}(Q)((V^{n+1})^{k} - V^{n+1}) \right\}.$$
(9.3)

If \hat{Q} satisfies

$$\hat{Q} \in \arg\max_{Q} \left\{ \mathcal{A}(Q)((V^{n+1})^{k+1} - V^{n+1}) + \mathcal{B}(Q)((V^{n+1})^k - V^{n+1}) \right\}. \tag{9.4}$$

then, from Eq. (9.3), we have, $(E^{k+1} = (V^{n+1})^{k+1} - V^{n+1})$

$$\mathcal{A}(\hat{Q})E^{k+1} \le \mathcal{B}(\hat{Q})E^k - \mathcal{E}^k \,, \tag{9.5}$$

or, since $\mathcal{A}(Q)$ is an M matrix,

$$E^{k+1} \leq \mathcal{A}(\hat{Q})^{-1} \mathcal{B}(\hat{Q}) E^k - \mathcal{A}(\hat{Q})^{-1} \mathcal{E}^k \leq C_1 \|E^k\|_{\infty} \mathbf{e} + C_2 \|\mathcal{E}^k\|_{\infty} \mathbf{e}$$

$$C_1 = \max_{Q} \|\mathcal{A}(Q)^{-1} \mathcal{B}(Q)\|_{\infty}$$

$$C_2 = \max_{Q} \|\mathcal{A}(Q)^{-1}\|_{\infty}$$

$$(9.6)$$

where $\mathbf{e} = [1, 1, \dots, 1]'$. Similarly

$$\mathcal{E}^{k} = \max_{Q} \left\{ -\mathcal{A}(Q)(V^{n+1})^{k+1} + \mathcal{B}(Q)(V^{n+1})^{k} + \mathcal{C}(Q, V^{n}) \right\} - \max_{Q'} \left\{ -\mathcal{A}(Q')V^{n+1} + \mathcal{B}(Q')V^{n+1} + \mathcal{C}(Q', V^{n}) \right\} \geq \min_{Q} \left\{ -\mathcal{A}(Q)((V^{n+1})^{k+1} - V^{n+1}) + \mathcal{B}(Q)((V^{n+1})^{k} - V^{n+1}) \right\}.$$
(9.7)

Hence if

$$\bar{Q} \in \arg\min_{Q \in Z} \left\{ -\mathcal{A}(Q)((V^{n+1})^{k+1} - V^{n+1}) + \mathcal{B}(Q)((V^{n+1})^k - V^{n+1}) \right\}, \quad (9.8)$$

then

$$E^{k+1} \ge \mathcal{A}(\bar{Q})^{-1} \mathcal{B}(\bar{Q}) E^k - \mathcal{A}(\hat{Q})^{-1} \mathcal{E}^k \ge -C_1 \|E^k\|_{\infty} \mathbf{e} - C_2 \|\mathcal{E}^k\|_{\infty} \mathbf{e} . \tag{9.9}$$

Equations (9.6) and (9.9) then give

$$||E^{k+1}||_{\infty} \le C_1 ||E^k||_{\infty} + C_2 ||\mathcal{E}_{\max}||_{\infty} ||\mathcal{E}_{\max}||_{\infty} = \max_{k} ||\mathcal{E}^k||_{\infty}.$$
(9.10)



From [21], we have that

$$C_1 \le \frac{\theta \hat{\lambda} \Delta \tau}{1 + \theta (r + \hat{\lambda}) \Delta \tau}; \qquad \hat{\lambda} = \max_j \lambda_j < 1,$$
 (9.11)

and C_2 bounded, hence

$$||E^{k+1}||_{\infty} \le C_1^{k+1} ||E^0||_{\infty} + C_2 \left(\frac{1 - C_1^{k+1}}{1 - C_1} \right) ||\mathcal{E}_{\max}||_{\infty}, \tag{9.12}$$

and in view of Eq. (9.11), we obtain

$$\lim_{k \to \infty} \|E^{k+1}\|_{\infty} \le \frac{C_2}{1 - C_1} \|\mathcal{E}_{\max}\|_{\infty}. \tag{9.13}$$

Manipulation of Line 4 in Algorithm 6.2 results in

$$\mathcal{A}(Q^m)(U^{m+1} - U^m) = \max_{Q} \left\{ -\mathcal{A}(Q)U^m + \mathcal{B}(Q)(V^{n+1})^k + \mathcal{C}(Q, V^n) \right\}$$
(9.14)

Recall that at convergence of Algorithm 6.2, we have

$$(V^{n+1})^{k+1} = U^{m+1}. (9.15)$$

When Algorithm 6.2 terminates, we have

$$\mathcal{E}^{k} = \max_{Q} \left\{ -\mathcal{A}(Q)(V^{n+1})^{k+1} + \mathcal{B}(Q)(V^{n+1})^{k} + \mathcal{C}(Q, V^{n}) \right\}, \tag{9.16}$$

then, from Eqs. (9.14) and (9.16) (with $U^{m+1} = (V^{n+1})^{k+1}$)

$$\mathcal{E}^{k} = \mathcal{A}(Q^{m})(U^{m+1} - U^{m}) + \max_{Q} \left\{ -\mathcal{A}(Q)U^{m+1} + \mathcal{B}(Q)(V^{n+1})^{k} + \mathcal{C}(Q, V^{n}) \right\}$$

$$- \max_{Q} \left\{ -\mathcal{A}(Q)U^{m} + \mathcal{B}(Q)(V^{n+1})^{k} + \mathcal{C}(Q, V^{n}) \right\}$$
(9.17)

and then

$$|\mathcal{E}^k| \le |\mathcal{A}(Q^m)(U^{m+1} - U^m)| + \max_{Q} |A(Q)(U^{m+1} - U^m)|, \qquad (9.18)$$

which implies

$$\|\mathcal{E}^k\|_{\infty} \le 2 \cdot \max_{Q} \|A(Q)\|_{\infty} \|U^{m+1} - U^m\|_{\infty}. \tag{9.19}$$

Since $\mathcal{A}(Q^m)$ is bounded, $\|\mathcal{E}^k\|_{\infty}$ (and hence $\|\mathcal{E}_{\max}\|_{\infty}$) can be made arbitrarily small by making *tolerance*_{inner} small in Algorithm 6.2.

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