

COMMENT ON “CONSTRAINED OPTIMIZATION APPROACHES TO ESTIMATION OF STRUCTURAL MODELS”

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We revisit the comparison of mathematical programming with equilibrium constraints (MPEC) and nested fixed point (NFXP) algorithms for estimating structural dynamic models by Su and Judd (2012). Their implementation of the nested fixed point algorithm used successive approximations to solve the inner fixed point problem (NFXP-SA). We redo their comparison using the more efficient version of NFXP proposed by Rust (1987), which combines successive approximations and Newton–Kantorovich iterations to solve the fixed point problem (NFXP-NK). We show that MPEC and NFXP are similar in speed and numerical performance when the more efficient NFXP-NK variant is used.

KEYWORDS: Structural estimation, dynamic discrete choice, NFXP, MPEC, successive approximations, Newton–Kantorovich algorithm.

1. INTRODUCTION

IN “CONSTRAINED OPTIMIZATION APPROACHES TO ESTIMATION OF STRUCTURAL MODELS,” Su and Judd (2012; hereafter SJ) proposed a constrained optimization approach for maximum likelihood estimation of infinite horizon dynamic discrete choice models—mathematical programming with equilibrium constraints (MPEC). They argued that MPEC is superior to the *nested fixed point* (NFXP) algorithm proposed by Rust (1987). NFXP uses the fact that the likelihood depends on the parameters via the *value function* to a dynamic programming (DP) problem. Under weak conditions, the value function is the unique fixed point to a contraction mapping defined by the *Bellman equation* to the DP problem and is a smooth implicit function of the underlying structural parameters of the problem. NFXP uses this to maximize the likelihood using standard *unconstrained* quasi-Newton optimization algorithms, except that each time the likelihood is evaluated, NFXP calls a fixed point subroutine to compute the value function corresponding to the current parameter values.

In contrast, the MPEC method does not need a specialized inner loop algorithm to compute the fixed point. Instead, it recasts the problem of maximizing

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the likelihood function as a constrained optimization problem with respect to the K structural parameters plus N additional variables, which are the values of value function at a set of N grid points in the state space. These N additional variables must satisfy the Bellman equation, which can be recast as N “side constraints.” Thus MPEC also implicitly solves the fixed point problem while searching for structural parameter values that maximize the likelihood, but using a general-purpose *constrained* optimization algorithm.²

SJ used the model of optimal replacement of bus engines of Rust (1987) to conduct a Monte Carlo study to compare the performance of the MPEC and NFXP algorithms. They found that MPEC outperformed NFXP in terms of CPU time by up to three orders of magnitude.

The point of this comment is to note that SJ used a version of NFXP we refer to as “NFXP-SA” where the method of *successive approximations* (SA) is used to solve the inner fixed point problem. However, it is well known that successive approximations is an inefficient algorithm for computing fixed points of contraction mappings, especially when the modulus of the contraction (which equals the discount factor in the underlying dynamic programming problem) is close to 1.

We redo the SJ Monte Carlo study using the more efficient version of NFXP that Rust (1987) employed. This version, NFXP-NK, combines successive approximations and Newton–Kantorovich (NK) iterations to solve the inner fixed point problem significantly faster and more reliably, especially when the discount factor is close to 1. We show that NFXP and MPEC are roughly equivalent in their numerical performance when the more efficient NFXP-NK variant is employed.

NK is the preferred method for computing contraction fixed points because it has guaranteed quadratic convergence rate in its domain of attraction. However, NK is only locally convergent, so the original design of the NFXP algorithm (Rust (1987, 2000)) starts with successive approximations to ensure global convergence, and switches to NK iterations only after it detects that the domain of attraction has been reached.³ This hybrid algorithm or “polyalgorithm” ensures that a highly accurate solution can be found after only a small combined number of iterations. In particular, the combination of these two approaches makes the performance of the NFXP-NK algorithm independent of

²The specific implementation SJ used is KNITRO (see Byrd, Nocedal, and Waltz (2006)), which is run under AMPL software that provides analytic first and second order derivatives, and under Matlab with analytic gradients and the Hessian approximated numerically.

³Rust (2000, p. 28) detailed the theory of when the algorithm has to switch from successive approximations to NK iterations. The idea is based on the fact that the ratio of tolerances in two successive iterations of SA algorithm approaches the modulus of contraction (β) as the algorithm progresses. When this ratio is close enough to β , the shape of the value function is nearly recovered (one can show that this ratio equals β exactly when the current iteration differs from the fixed point by a constant λ), which is a signification that the domain of attraction of the NK method has been reached (λ will be “stripped away” in just a single iteration).

the value of the discount factor β , whereas the CPU times of the NFXP-SA algorithm steadily increase as $\beta \rightarrow 1$, as shown in Table II of SJ (p. 2228).

2. RESULTS

Table I presents a comparison of CPU times for MPEC and NFXP that reproduces Table II of SJ. While we have developed our own code to implement NFXP-NK, we have been using the same setup and computer code for MPEC that was used to produce the results of the original paper by SJ.⁴ We have replicated SJ's results for NFXP-SA, but since this version of NFXP is inefficient and is completely dominated by NFXP-NK, we only report results for the latter in Table I. Similarly, we skip the results regarding MPEC-Matlab implementation which uses first order analytical derivatives only, because its run times were about two orders of magnitude larger than those of MPEC-AMPL

TABLE I
MPEC VERSUS NFXP-NK: SAMPLE SIZE 6000^a

β	Converged (Out of 1250)	CPU Time (in Sec.)	# of Major Iter.	# of Func. Eval.	# of Bellm. Iter.	# of N-K Iter.
MPEC-AMPL						
0.975	1246	0.054	9.3	12.1		
0.985	1217	0.078	16.1	44.1		
0.995	1206	0.080	17.4	49.3		
0.999	1248	0.055	9.9	12.6		
0.9995	1250	0.056	9.9	11.2		
0.9999	1249	0.060	11.1	13.1		
NFXP-NK						
0.975	1250	0.068	11.4	13.9	155.7	51.3
0.985	1250	0.066	10.5	12.9	146.7	50.9
0.995	1250	0.069	9.9	12.6	145.5	55.1
0.999	1250	0.069	9.4	12.5	141.9	57.1
0.9995	1250	0.078	9.4	12.5	142.6	57.5
0.9999	1250	0.070	9.4	12.6	142.4	57.7

^aThis table is a replication of Table II in Su and Judd (2012) with NFXP-SA replaced by NFXP-NK (section on MPEC-Matlab is skipped to conserve space). For each β , five starting points were used for each of the 250 simulated samples. CPU time, number of major iterations, number of function evaluations, and number of inner loop iterations are the averages over the convergent runs. Inner loop iterations include both value function iterations and Newton-Kantorovich iterations.

⁴SJ have kindly provided well documented MPEC code for this problem via Che-Lin Su's website. We are thankful to Che-Lin Su for pointing out numerical instability of the original code due to the lack of recentering of the value functions in accordance with Rust (2000, p. 27). Recentering has been implemented in the numerically stable version of MPEC code we use here. Our NFXP code is available on request.

in all our experiments.^{5,6} To conserve space, we also omit the table of structural estimates (Table I of SJ), which shows that, conditional on convergence, both methods were able to recover the structural parameters of the model. It is also the case in all our experiments.

It is clear from Table I that when the fast NFXP-NK algorithm is used, the CPU times are comparable to those of MPEC-AMPL, the implementation utilizing first and second order analytic derivatives of the objective function. It takes both methods about 0.07 to 0.08 of a second to structurally estimate the bus engine replacement model. Also, unlike SJ, we find NFXP-NK more reliable than MPEC-AMPL, as indicated by the second column in Table I.

As the second panel of Table I shows, NFXP-NK uses far fewer successive approximation steps compared to the NFXP-SA results in Table II of SJ, and this is the main reason why NFXP-NK is so much faster. For the highest discount factor $\beta = 0.995$ in Table II of SJ, an average of nearly 750,000 successive approximation iterations were required, compared to just 145 for NFXP-NK. With an average of 55 NK iterations per estimation, the average number of both types of inner loop iterations for NFXP-NK is remarkably insensitive to the discount factor. On average, it takes NFXP-NK only 12 successive approximation steps and 4 NK iterations per function evaluation to compute a highly accurate fixed point (to a tolerance of 10^{-13}) when $\beta \geq 0.9995$.

3. CONCLUSION

Our findings lead us to a different conclusion from SJ, namely, that “Monte Carlo results confirmed that MPEC is significantly faster than NFXP, particularly when the discount factor in the dynamic-programming model is close to 1.” (p. 2228). We have shown that this conclusion is an artifact of their use of an inefficient version of NFXP, NFXP-SA, which uses successive approximations to solve the fixed point problem. When we compare MPEC to the more efficient implementation of NFXP that Rust (1987) originally proposed,

⁵We used the same specification of the bus engine replacement model, including the same true parameter values, same sample size and fixed point dimension, and the same number of Monte Carlo replications as in Table II of SJ. We fixed the stopping tolerance for the inner fixed point in the NFXP-NK algorithm at 10^{-13} as in Rust (1987), which is 1/1000 of the stopping tolerance 10^{-10} that SJ used for NFXP-SA. Similarly to SJ, we estimated transition probabilities for mileage travelled by buses jointly with other parameters (replacement cost and maintenance cost) by maximizing the full likelihood function, following the partial likelihood optimization as described in Rust (1987, 2000). We used the BHHH algorithm on the outer loop of NFXP-NK and frequency based starting values for the transition probability parameters for both methods. For the polyalgorithm, we use a minimum of $minstp = 2$ and maximum $maxstp = 20$ successive approximation iterations and a relative tolerance to switch from SA to NK iteration $rol = 0.02$; see Rust (2000, p. 28).

⁶Our hardware and software setup was the following: Mac Book Pro with 2.3 GHz Intel Core i7 processor and 8 GB of memory with OS X version 10.9.5, Matlab 2014a constrained to a single core with `-singleCompThread` startup option, AMPL Version 20131213 and KNITRO 9.0.1.

NFXP-NK, we find that NFXP and MPEC are approximately equally fast and accurate.

There is a fundamental difference between how NFXP and MPEC solve the structural estimation problem. In the case of NFXP, the choice probabilities entering the likelihood function are computed independently of the data in the inner loop. For MPEC, both the fixed point calculation and the maximization of the likelihood are done simultaneously, which not only implies that the gradient vector and the Hessian matrix are both high dimensional objects, but also that the whole data set needs to be processed multiple times when computing nonzero elements of these objects. The separation between solving the model and computing the likelihood enables NFXP to use traditional unconstrained quasi-Newton/gradient search algorithms for likelihood maximization—such as the Berndt–Hall–Hall–Hausman (BHHH) algorithm (Berndt, Hall, Hall, and Hausman (1974))—over a relatively small number of structural parameters. Unlike MPEC, NFXP recognizes the fact that the objective function is a sum of individual likelihoods each of which is computed from the set of value functions that are smooth in the structural parameters. The BHHH algorithm exploits the information identity to approximate the Hessian of the likelihood with the negative of the outer product of the scores. Therefore, because the Hessian approximation is always negative semi-definite, BHHH always moves in the direction of the gradient (i.e., towards the maximum), even in convex areas of the likelihood function. Hence, beyond the advantage of avoiding computation of second order derivatives, BHHH has the major advantage of always moving uphill for small enough step size, and thus is globally convergent to at least local maximum of the likelihood function. The robustness and computational efficiency of NFXP comes from fully exploiting the structure of the maximum likelihood estimation problem, that is, by recognizing that the Bellman operator is a contraction mapping and that objective function is a sample sum over individual likelihoods.

We believe that MPEC has many desirable features, the most important of which is ease of use by people who are not interested in devoting time to the special-purpose programming necessary to implement NFXP-NK. Our results indicate that MPEC is very fast and competitive with NFXP-NK in the bus engine replacement model, and, particularly in conjunction with intuitive AMPL language, it could save many users substantial programming time and enable them to structurally estimate many models of interest. For this reason, MPEC may be the method of choice for structural estimation of relatively well behaved infinite horizon models that can be formulated using software such as AMPL.

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