## Structural change and the dynamics of energy prices \*

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#### **ABSTRACT**

We test for the presence of time-varying parameters in long-run real energy prices as suggested in the class of models proposed by Pindyck (1999). These models postulate mean-reverting prices with continuous and random changes in their level and trend, and are estimated using Kalman filtering techniques. Since the latter contain boundary as well as possibly unidentified parameters when there is no parameter variation, the test statistics have non-standard distributions and involve nuisance parameters. To solve the statistical difficulties associated with this problem, we use a simulation-based method, called *maximized Monte Carlo* tests [Dufour (2002)], which can yield provably exact tests for highly irregular problems even in small samples. Significant parameter variation is detected for both oil and natural gas prices, but not for coal price. Out-of-sample forecasts are also calculated to differentiate between alternative models. Contrary to Slade (1982) who uses a deterministic trend model, we find no evidence of U-shaped trend for oil or coal prices; there is almost no upward trend either. In contrast, the natural gas price trend has an upward U-shaped form.

**Keywords:** structural change; energy prices; coal; gas; crude oil; mean reversion; exact test; Monte Carlo test; Kalman filter.

Journal of Economic Literature classification: C22, C52, C53, Q40.

#### **RÉSUMÉ**

Nous testons la présence de paramètres variables dans les équations suggérées par Pindyck (1999) afin de modéliser les prix à long terme de l'énergie. Ces modèles postulent des processus de retour à la moyenne avec des changements continus et aléatoires des coefficients de la tendance, et sont estimés en utilisant des techniques de filtrage de Kalman. Comme ces derniers comprennent des paramètres sur la frontière d'admissibilité ou non-identifiés en l'absence de changement structurel, les distributions des statistiques de test correspondantes sont typiquement non-standards et dépendent de paramètres de nuisance. Pour résoudre les difficultés statistiques associées à ces problèmes, nous appliquons une procédure basée sur des simulations, la méthode des tests de Monte Carlo maximisés [Dufour (2002)], laquelle fournit des tests dont le niveau est contrôlé pour des problèmes hautement irréguliers, même sur de petits échantillons. Nous détectons des paramètres variables dans le cas du pétrole brut et du gaz naturel, mais pas pour le charbon. Afin de comparer divers modèles qui semblent acceptables par des critères de tests, nous calculons des prévisions hors-échantillon.

**Mots clés :** changement structurel ; prix de l'énergie; charbon; gaz naturel; pétrole brut; retour à la moyenne; test exact; test de Monte Carlo; test de Monte Carlo maximisé; bootstrap; filtre de Kalman.

Classification du Journal of Economic Literature : C22, C52, C53, Q40.

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## 1. Introduction

It is generally recognized that fluctuations in energy prices have important and lasting effects on the economies of industrialized countries. As a recent example, Hamilton (2003) found a strongly significant nonlinear relationship between changes in oil prices and GDP growth. Similarly, in small open-economy settings, Amano and van Norden (1998) found long-run links between oil prices and real exchange rates. Enduring price movements in energy commodities can also cause relative price changes among a wide range of products in the economy, which can then influence the rate of inflation over an extended period.<sup>1</sup>

Interpreting and predicting the behavior of energy prices have proved to be quite challenging. In addition to domestic and international supply and demand conditions, a complete model also needs to take into account market regulations, technological advances, and geopolitical considerations. These non-market-related aspects present the biggest challenges. For that reason, a fully articulated structural model may be quite difficult to build and unreliable, and Pindyck (1999) has suggested that a more practical approach for long-run forecasting may consist in building simple flexible dynamic models which incorporate implicitly the role played by demand shifts, technological change and depletion. These models are more general than models with deterministic trends as used by Slade (1982) to analyze the long term behavior of depletable resource prices. Although the latter models are special cases of the general ones under boundary parameter restrictions, standard statistical tests can be misleading because of boundary nesting and the application of appropriate tests is a contribution of this paper.

We reconsider the class of trend models with time varying parameters (TVP) proposed by Pindyck (1999) for that purpose. Despite their simplicity, these models are quite flexible, allowing both random walks with drift and/or changing trend lines where prices revert to a possibly moving mean. An important feature of the proposed models is the inclusion of time varying trend parameters. Different forms of the model may reflect different assumptions on resource depletion and technological change. For example, a form of energy which is produced and sold competitively should have its price reverting to a long-run marginal cost which itself can change over time. Using a simple Hotelling model, Pindyck showed that long-run energy prices should revert to an *unobservable* trending long-run marginal cost, with continuous random changes in the level and slope of the trend. A family of econometric models for these prices that integrate desired features (discrete versions of multivariate Ornstein-Uhlenbeck processes) was also suggested for energy prices. Alternative versions of these models were estimated using the Kalman filter techniques, and out-of-sample forecasts were computed. The forecast exercises conducted by Pindyck contain some mixed results, but overall the class of models considered appears to be quite promising.

Pindyck (1999) did not, however, provide statistical tests for the class of models proposed. In particular, the time varying parameter specification was not tested statistically. Yet, the decision to use a TVP model (as opposed to a more standard autoregressive or fixed coefficient trend model) may have non-negligible consequences on model forecasts.

An important difficulty one meets here comes from the fact that TVP models contain boundary

<sup>&</sup>lt;sup>1</sup>There is no unanimity about the interpretation of the recent history in this respect. For a critical review, see Barsky and Kilian (2004).

as well as unidentified parameters, especially when there is no parameter variation, so that test statistics have non-standard distributions and involve nuisance parameters. Usual chi-square critical points can easily lead to spurious rejections even with fairly large data sets, because the regularity conditions underlying classical asymptotics fail.<sup>2</sup> Another problem is the fact that likelihood functions tend to be *ill-behaved* for empirically relevant parameter values. Thus, although sophisticated numerical recipes and global maximizers are readily available, it is well known that perfect maximization is not granted in this context.

Here, we complement Pindyck's (1999) study by conducting tests for continuous and random shifts in real energy prices. We are thus able to select, within the suggested family of models, specifications that are statistically justified for crude oil, coal, and natural gas prices. Our methodology relies on exact simulation-based test procedures, applicable even with small samples to highly irregular problems for which standard techniques are not valid. We also complement our results by providing a number of out-of-sample forecast statistics.

The merit of simulation-based procedures was recently demonstrated in a related framework, namely for the presence of jumps in the context of jump diffusions; see Dufour and Khalaf (2001), Khalaf, Saphores and Bilodeau (2003), and Saphores, Khalaf and Pelletier (2002). Those studies circumvent the unidentified nuisance-parameter problem through bounds Monte Carlo tests. These involve simulation-based cut-off points (*i.e.*, *p*-values) based on conservative bounds. The approach used in the present paper is related to the latter in the sense that it is also bounds-based. However, here, an *optimal* (*i.e.*, tightest) bound is provided. Specifically, we apply the *maximized Monte Carlo* (MMC) test technique [Dufour (2002)], which is based on comparing the maximal *p*-value of the test (obtained by simulation, over relevant nuisance parameters) with the significance level. Consequently, (i) level control is ensured by construction, and (ii) the tightest cut-off point is obtained numerically.<sup>3</sup>

Our findings indicate significant TVP effects in two of the three energy-price series examined, which supports Pindyck's proposed class of models. Indeed, we find significant parameter variation in the natural gas and oil series. Our results suggest a slow reverting process of log price to a fast adjusting random mean and a very slow adjusting trend. We find no evidence of U-shaped trend with the logs of oil and coal prices. There is almost no upward trend over 120 years. Forecast statistics also suggest that the TVP specification outperforms fixed-coefficient mean reverting models and random-walk-with-drift models, for all three price series.

In Section 2, we describe the class of proposed models and the test method used. Section 3 documents and discusses our empirical results. We conclude in Section 4.

### 2. Model and test method

Pindyck (1999) considers a basic Hotelling model for a depletable resource produced in a compet-

<sup>&</sup>lt;sup>2</sup>See Hansen (1996), Dufour (1997, 2003) and Andrews (2000, 2001).

<sup>&</sup>lt;sup>3</sup>For further discussion of Monte Carlo test method in econometrics, see, for example, Dufour and Kiviet (1996, 1998), Kiviet and Dufour (1997), Dufour, Farhat, Gardiol and Khalaf (1998), Dufour and Khalaf (2001, 2002a, 2002b), Khalaf and Kichian (2002, 2003, 2004), Dufour, Khalaf, Bernard and Genest (2004), and Dufour and Jouini (2005).

itive market. With constant marginal cost of extraction c and an isoelastic demand function with unitary elasticity, the price level is given by

$$P_t = c + [(ce^{rt}/(e^{rcR_0/A} - 1))]$$
(2.1)

where  $R_0$  is the initial stock of the depletable resource, A is a demand shifter, and r is the interest rate. This implies that the slope of the price trajectory is given by

$$dP_t/dt = rce^{rt}/(e^{rcR_0/A} - 1),$$
 (2.2)

so that changes in demand, extraction costs, and reserves all affect this slope. For example, an increase in A causes the slope to increase, while increases in c or  $R_0$  reduce the slope. In addition, increases in c or A cause the price level to increase, whereas an increase in  $R_0$  leads to a decrease in this level. If, as Pindyck (1999) argues, these factors fluctuate in a continuous and unpredictable manner over time, then long-run energy prices should revert to a trend which itself fluctuates in the same fashion.

A class of models that integrates the above features is the generalized Ornstein-Uhlenbeck process. Pindyck (1999) proposes a discretized version of this model as a suitable econometric framework for forecasting long-run energy prices. This leads to the following AR(1)-type dynamic model:

$$P_t = c_1 + \phi_{1t} + \phi_{2t}t + c_2P_{t-1} + \epsilon_t, \quad t = 1, \dots, T,$$
(2.3)

where  $P_t$  refers to the log real price of an energy product and the coefficients  $\phi_{1t}$  and  $\phi_{2t}$ , follow the stochastic processes

$$\phi_{1t} = c_3 \phi_{1,t-1} + v_{1t} \,, \tag{2.4}$$

$$\phi_{2t} = c_4 \phi_{2t-1} + v_{2t}. \tag{2.5}$$

The processes for  $\phi_{1t}$  and  $\phi_{2t}$  are unobservable, continuously evolving parameters which reflect long-run marginal costs including scarcity rent, in the underlying structural model.

For tractability purposes, we estimate and test this model separately for each price series considered. Clearly, a multivariate analysis (*i.e.* joint estimation) may be preferable, at least in principle, since energy sources are substitutes in the long run. Yet given sample size limitations, numerical burdens (pointed out by Pindyck) may outweigh efficiency gains for this particular application. Further, modelling jointly all the equations involves additional specification choices that may be disputable.

For similar considerations, we also impose the following distributional assumptions on the univariate process (2.3). The underlying error terms  $\epsilon_t$ ,  $v_{1t}$ , and  $v_{2t}$ ,  $t=1,\ldots,T$ , are assumed to be independently and identically normally distributed with zero means and covariances, and variances  $\sigma_{\epsilon}^2$ ,  $\sigma_{v_1}^2$ , and  $\sigma_{v_2}^2$ , respectively. Stationarity constraints (on  $c_2$ ,  $c_3$  and  $c_4$ ) are (at least in principle) not necessary. It is worth noting that despite the latter simplifying hypotheses, the statistical

framework remains non-regular, as we will explain in what follows.

Assuming normality of  $\epsilon_t$ ,  $v_{1t}$ , and  $v_{2t}$ , Pindyck proposes that Kalman filtering be applied to obtain paths for the state variables  $\phi_{1t}$  and  $\phi_{2t}$ . This means that, starting with initial values for model parameters and state variables, the filter computes at each period new values for the state variables to reflect new information on the observable series. Once the full paths of the state variables are determined, the model can be estimated by maximum likelihood to obtain parameter estimates. For the model above, details of the Kalman filtering procedure are described in Appendix B. In view of assessing the statistical significance of TVP effects, the null hypothesis of interest is a simple mean-reverting model around a fixed trend line [the trending Ornstein-Uhlenbeck process given by equation (24) in Pindyck (1999)], i.e.,

$$P_t = c_1 + \phi_1 + \phi_2 t + c_2 P_{t-1} + \epsilon_t, \quad t = 1, \dots, T.$$
 (2.6)

It is clear that the models to be compared statistically are nested at the boundaries of certain parameters; formally,

model (2.6) 
$$\subseteq$$
 model (2.3) when  $\sigma_{v_1}^2 \to 0, \ \sigma_{v_2}^2 \to 0, \ \text{and} \ c_3 = c_4 = 1$ . (2.7)

Further, it is easy to see that some parameters may not be identifiable under certain parameter configurations: for example, the constant term is not identified when  $c_3 = 1$  and  $\sigma_{v_1}^2 = 0$  and it is "poorly" identified when we are close to these values.

In this context, one cannot rely on estimated standard errors and standard limiting distributions, since their use for building tests and confidence sets is not justified even asymptotically. In particular, the distributions of some widely used test statistics, such as t-type and more generally Wald-type statistics, may be difficult (if not impossible) to bound under various null hypotheses of interest, so that controlling the level of such tests can be quite difficult. By contrast, the distributions of likelihood-ratio-type statistics appear to be more stable and thus provide a more appropriate basis for statistical inference; see Dufour (1997, 2003) and Stock, Wright and Yogo (2002).

Taking into account these observations and the potential computational cost of maximum likelihood (ML) estimation of TVP models, we focus here on a quasi-likelihood-ratio (QLR) type statistic of the form:

$$QLR(J) = 2[L_{TVP}(J) - L_{FCM}]$$
(2.8)

where  $L_{TVP}(J)$  and  $L_{FCM}$  are, respectively, the maximum of the log-likelihood functions associated with (2.3) and (2.6), J is the maximum number of iterations allowed in the numerical maximization exercise under (2.3), and the subscript FCM stands for fixed-coefficient model. J is explicitly spelled out to take into account the fact that the number of iterations has an influence on the finite-sample distribution of the test statistic: depending on the number of iterations (along with a convergence criterion), one really considers different test statistics.

In view of the boundary and identification difficulties mentioned above, it would be wrong to

<sup>&</sup>lt;sup>4</sup>For further discussion of this issue, see Robinson (1988).

compare the QLR statistic with standard tabulated values from a  $\chi^2$  table. Both the finite-sample and asymptotic distributions of the QLR statistic may be strongly affected by the irregularities inherent to the problem studied here.<sup>5</sup> We need a procedure that can take these features into consideration.

To test the hypotheses of interest in this context, we therefore resort to maximized Monte Carlo (MMC) tests [Dufour (2002)], which we apply to the QLR statistic. Let us denote the vector of nuisance parameters that appear under the null hypothesis as:

$$\theta = \{\lambda, \, \phi_2, \, c_2, \, \sigma_{\epsilon}^2\}, \, \lambda = c_1 + \phi_1. \tag{2.9}$$

In Appendix A, we provide a more detailed exposition of the MMC test method for a general test statistic whose null distribution can be simulated given a vector of nuisance parameters  $\theta$ . Herein, we will summarize the technique as it applies to our testing problem, where  $\theta$  is given by (2.9). In our test procedure, all that is needed to obtain draws from the null data-generating process is to set a value for  $\theta$ ; the unidentified nuisance parameters (for example,  $c_3$  and  $c_4$  under the constant coefficient model) simply do not appear in the distribution. More precisely, we proceed as follows.

- (i) We calculate the likelihood ratio statistic (2.8) using the maximized likelihood value of the TVP model (2.3) the *alternative model* and the one of the constant-coefficient model (2.6) the null model. In the process, we save the quasi-maximum-likelihood estimate of  $\theta$  imposing (2.6) and the observed value of QLR(J). We denote these values  $\hat{\theta}_{FCM}$  and  $QLR_0(J)$  respectively.
- (ii) We generate data from the null model drawing from the normal distribution, setting  $\theta$  equal to its estimated value  $\hat{\theta}_{FCM}$ , reestimate the restricted and the alternative models from these simulated data, and compute the corresponding  $QLR(J; \hat{\theta}_{FCM})$  where the presence of  $\hat{\theta}_{FCM}$  indicates that  $QLR(J; \hat{\theta}_{FCM})$  depends on data simulated after setting  $\theta = \hat{\theta}_{FCM}$ . This process is repeated N times, yielding N simulated test statistics from a data generating process (DGP) that satisfies the null hypothesis:  $QLR_i(J; \hat{\theta}_{FCM}), i = 1, \ldots, N$ .
- (iii)  $QLR_0(J)$  is compared with this distribution and a p-value is calculated based on the rank of  $QLR_0(J)$  relative to its simulated counterparts; see equations (A.2), (A.3), and (A.4) in Appendix A. We will call the number so obtained the *local Monte Carlo* (LMC) p-value.

The MMC technique involves repeating step (ii) above, sweeping over combinations of admissible values of  $\theta$  instead of  $\hat{\theta}_{FCM}$ . This can be viewed as a Monte Carlo implementation of the standard definition of the level of a test in the presence of nuisance parameters: when a test is nuisance-parameter dependent, an  $\alpha$  level is achieved by comparing the largest p-value over all nuisance parameters consistent with the null hypothesis to  $\alpha$  [see Lehmann (1986)]. The MMC method, by construction, works exactly in this way. Thus, we obtain a MC p-value for each value of  $\theta$ . The MMC p-value is then the highest MC p-value so obtained. Since the maximized p-value function is a non-differentiable step function, we use simulated annealing (a global non-gradient-based algorithm) to obtain the maxima [see Goffe, Ferrier and Rogers (1994)]. This provides, by

<sup>&</sup>lt;sup>5</sup>See Andrews (2000, 2001), Dufour (1997, 2003) and Stock et al. (2002).

construction, the tightest bound p-value. The MMC test is significant at level  $\alpha$  if the MMC p-value (say  $p_{MMC}$ ) is less than or equal to  $\alpha$ :

$$p_{MMC} \le \alpha. (2.10)$$

Of course, if the MC p-value obtained in step (ii) with  $\theta = \hat{\theta}_{FCM}$  exceeds  $\alpha$  (e.g., 10 per cent), there is no need to proceed with the maximization; this saves execution time. Test results are reported in Section 3.

If the MC *p*-value is computed using a single (consistent) parameter estimate of model nuisance parameters, we get a *local MC* (LMC) test or, equivalently, a parametric bootstrap test. Bootstrap procedures tend to be considerably more reliable than procedures based on asymptotic critical values. In the context of our problem, however, where the asymptotic distribution may depend in a discontinuous way on nuisance parameters, it is well known that bootstrap procedures may also fail even asymptotically.<sup>6</sup> By contrast, the MMC procedure is immune to such failures [see Dufour (2002)]. In the following section, we report LMC tests along with MMC test results.

## 3. Empirical results

We consider the annual data set analyzed by Pindyck (1999).<sup>7</sup> The series for crude oil and bituminous coal extend from 1870 to 1996, while, for natural gas, the data cover 1919 to 1996. The nominal oil price series and the natural gas series for 1870 to 1973 are from Manthy (1978) and the U.S. Bureau of the Census (1975). Pindyck (1999) updated this series through 1995 using data from the U.S. Energy Information Agency and, for 1996, the *Wall Street Journal*. The series are deflated using the U.S. wholesale price index until 1970, and the producers price index thereafter. Estimation is conducted on the logarithm of real prices.

First, we test the constant coefficient model against Pindyck's general TVP specification for each energy product. In other words, we compute the QLR statistic for testing the fixed coefficient model (2.6) against the TVP model (2.3) imposing only stability restrictions:

$$H_{TVP}: |c_2| < 1, 0 < c_3 < 1, 0 < c_4 < 1.$$
 (3.1)

Such restrictions as in (3.1) are not necessary for the validity of our test procedure. Yet, for this particular application, we have observed that stability constrains did enhance convergence and avoid corner solutions. Maximization is typically difficult to achieve in TVP contexts, and the numerical burden tends to be relatively heavy. Indeed, to obtain the MC *p*-values [see step (ii) in the previous section], we run numerical nonlinear optimization algorithms 200 times [once with observed data (to derive the observed tests statistic) and 199 times with simulated data (to derive the simulated

<sup>&</sup>lt;sup>6</sup>See Athreya (1987), Basawa, Mallik, McCormick, Reeves and Taylor (1991), Sriram (1994), Andrews (2000), Inoue and Kilian (2002, 2003).

<sup>&</sup>lt;sup>7</sup>The data were generously provided by Pindyck.

Table 1. Tests of constant parameter model and Pindyck's specifications

Model tested $(H_0)$ : $P_t = c_1 + \phi_1 + \phi_2 t + c_2 P_{t-1} + \epsilon_t$							
Alternative model $(H_1)$	Energy type	LR	LMC	MMC			
$P_t = c_1 + c_2 P_{t-1} + \phi_{1t} + \phi_{2t} t + \epsilon_t$	Oil	14.7973	.005	.010			
$\phi_{1t} = c_3 \phi_{1t-1} + v_{1t}$	Coal	2.5973	.195	-			
$\phi_{2t} = c_4 \phi_{2t-1} + v_{2t}$	Gas	35.0205	.005	.005			

Note – The table reports p-values. LMC refers to the local MC (or parametric bootstrap) p-value and MMC to the maximized Monte Carlo p-value. The number of replications used for the MC tests is N=199. Since the MMC p-value must be larger than the LMC p-value, it is not necessary to report the MMC p-value if the former is larger than the level of the test (in this case, 5%).

test statistics)] for each admissible values of  $\theta$ . Furthermore, the underlying simulated samples are generated imposing the null (constant coefficient) model; so in this case, the maximization problem is by construction, ill conditioned.

To circumvent these difficulties, we applied the following numerical improvements. First, optimum results obtained by gradient-based algorithms are validated using global maximizers. For that purpose, we used *simulated annealing* [see Goffe et al. (1994)], particularly with the observed samples. Second, following a "normal convergence" output, the associated QLR statistics are checked for positivity. Indeed, if a negative QLR is obtained which signals non-convergence, the maximization algorithm is re-initiated using the "imperfect" solution as a starting value, until positivity is secured.

The results obtained are reported in Table 1, where LMC refers to local MC (parametric bootstrap) p-values and MMC to maximized p-values.<sup>8</sup> The maximal number of iterations allowed before the estimation process stops is  $J=120.^9$  All the MC tests are implemented using N=199 replications and the algorithm that maximizes the p-value function (in terms of  $\theta$ ) is initialized at the value used for the LMC test.

We see from the results in Table 1 that the constant coefficient model is rejected (by MMC tests) against the general TVP model for gas and oil prices at conventional significance levels, but not for coal prices (by the LMC procedure, hence also by the MMC procedure). Parameter estimates are reported in Table 2. We observe a slow adjustment process for all price series (estimates of  $c_2$  exceed .5) which reverts to a fast adjusting random mean (estimates of  $c_3$  are less than .5) and a random trend which has a very slow adjustment coefficients particularly for gas (estimates of  $c_4$  exceed .8). These results indicate good overall statistical support for Pindyck's proposed class of models: there are significant TVP effects in two of the three price series examined. Our test conclusions may also

<sup>&</sup>lt;sup>8</sup>For the regression coefficients, the values of  $\theta$  which are explored are determined by taking 5 standard errors on both sides of each component  $\theta_i$  of  $\theta$  ( $\hat{\theta}_{iFCM} \pm 5SE_i$ ). This is indeed quite wide (10 standard errors). Given that the estimators of the constant coefficient model converge at a rate at least as large as  $T^{1/2}$ , the box so obtained can be interpreted as a "consistent set estimator" of the relevant parameters [see Dufour (2002)]. For the error variances, we used the fixed interval [0.0001, 2], where the upper bound is well over all the estimates of these parameters.

<sup>&</sup>lt;sup>9</sup>Typically, in the examples considered here, convergence was achieved with much less than 80 iterations.

Table 2. Parameter estimates for the considered energy price models

Model	Туре	$c_1$	$c_3$	$c_2$	$c_4$	$\sigma_{v_1}$	$\sigma_{v_2}$	$\sigma_{arepsilon}$
$P_{t} = c_{1} + c_{2}P_{t-1} + \phi_{1t} + \phi_{2t}t + \epsilon_{t}$	Oil	.3508	.4799	.7415	.8898	6.3E-5	.0206	.4031
$\phi_{1t} = c_3 \phi_{1t-1} + v_{1t}  \phi_{2t} = c_4 \phi_{2t-1} + v_{2t}$	Coal Gas	.3437 2.7144	.1150 .0833	.8004 .5247	.9297 .9613	.2814 .1766	.0140 .0383	.0193 .0080
Model			ı	$c_2$	$\phi_2$		$\sigma_{arepsilon}$	
$P_t = \lambda + \phi_2 t + c_2 P_{t-1} + \epsilon_t$	Oil	.1894		.7869	.0016		.1944	
$\lambda = c_1 + \phi_1$	Coal	.12	88	.8976	.0	006	.08	861
,	Gas	.00	35	.9717	.0	011	.11	.94
Model	Type	λ		$c_2$ $\phi_2$		$\phi_2$	$\sigma_arepsilon$	
$P_t = \lambda + \phi_2 t + c_2 P_{t-1} + \epsilon_t$	Oil	02	95	1.0	.0	004	.20	75
$\lambda = c_1 + \phi_1$	Coal	.00	68	1.0	0	001	.08	377
$c_2 = 1$	Gas	02	285	1.0	.0	006	.11	95

explain why Pindyck's estimation results signal a poor fit with the coal price series.

Figure 1 shows our estimated price trends [formally, the estimated  $\phi_{2t}t$  term from (2.3)] over the sample period. The trend functions spike at the wake of the late seventies fuel-price shocks, yet following their impact, all three revert (rather slowly) to their respective long run mean. Interestingly, the latter is flat for both oil and coal prices, and slopes upward for natural gas. These results may be contrasted with Slade (1982)'s U-shaped curves, except for gas where in our case a U-shaped form is visible albeit less markedly than in Slade (1982). Recall however that the series analyzed by Slade ends in 1978; differing results may thus be driven by the post 1985 data.

One may note that estimates reported by Pindyck (1999, Table 2) signal "corner solution" values (duly discussed in the paper); the estimated  $c_3 = 1.0009$  for oil;  $c_4$  is imposed = 1 for coal, and the time-varying drift is omitted for gas in which case the estimated  $c_4$  is close to unity. Since we focus on (2.3) imposing (3.1) for all price series, our estimation results differ numerically from the ones reported by Pindyck. Differences may also reflect the sensitivity of Kalman filter estimates to filter initializations (a point raised by Pindyck). The question of forecast adequacy is thus worth re-visiting, to complement our analysis. So we produce forecasts as in Pindyck by estimating the TVP models for the samples ending in 1970 and 1980 respectively, and computing (continuously updated one step-ahead) dynamic forecasts out to the end of the full sample (*i.e.*, 1996). We then calculate the mean square errors (MSE's) for each of these forecast series and report them in Table 3. For comparison purposes, we also forecast MSE's obtained from a fixed-coefficient model and a unit root model, both estimated over samples ending in 1970 and 1980.

The out-of-sample one-step-ahead prediction errors are obtained as follows: given a sample of size T+j, we first remove j observations at the end of the sample that correspond to the forecast horizon considered. The model is then estimated on the remaining sample (i.e., until T), and the dependent variable's value is forecast for period T+1. The T+1 forecast error resulting from the comparison of the observed value of this variable and its estimated one is also computed. Next,

Table 3. Mean square forecast errors

Forecasting model	Energy type	MSE1971	MSE1981	
$P_t = \lambda + c_2 P_{t-1} + \phi_2 t + \epsilon_t$	Oil	0.378	0.086	
	Coal	0.099	0.108	
	Gas	1.676	3.096	
$P_t = \lambda + P_{t-1} + \phi_2 t + \epsilon_t$	Oil	0.206	1.241	
	Coal	0.105	0.323	
	Gas	1.161	1.005	
$P_{t} = c_{1} + c_{2}P_{t-1} + \phi_{1t} + \phi_{2t}t + \epsilon_{t}$ $\phi_{1t} = c_{3}\phi_{1t-1} + v_{1t}$ $\phi_{2t} = c_{4}\phi_{2t-1} + v_{2t}$	Oil	0.063	0.075	
	Coal	0.013	0.002	
	Gas	0.020	0.027	

Note – Mean square errors are calculated over 1971–96 and 1981–96, and denoted MSE1971 and MSE1981, respectively. The models in the first two rows are estimated by ordinary least squares.

the T+1 observed value of the dependent variable is added to our sample, and the model is reestimated. Now the T+2 observation is forecast, and the T+2 forecast error computed, and so on, and so forth, until all j observations are covered. This exercise allows parameter estimates and state variables to adjust to new information after 1970 and 1980. The results shown in Table 3 indicate that, over all forecast horizons, the TVP models yield superior forecasts for all series, including coal, although the TVP effects were not statistically significant for this series (the test p-value is around 20%). This holds true even when a unit-root specification is considered as an alternative model. The case of coal illustrates a well known challenge in time series analysis: the best fit model does necessarily lead to the best out-of-sample forecasts. Our results with this series relative to Pindyck's may also be explained by our continuously updated one-step ahead forecasting method.

Our comparative analysis underscores the merit of this model, and motivate further improvements, for example via multivariate approaches. Such specifications may also prove to be quite useful to model long-run components of two-factor short-run long-run models for resource prices.<sup>11</sup>

## 4. Conclusion

This paper has tested the statistical significance of Pindyck's (1999) suggested class of econometric equations that model the behavior of long-run real energy prices. The models postulate mean-reverting prices with continuous and random changes in their level and trend, and are estimated using Kalman filtering. In such contexts, the distributions of the test statistics are typically non-

 $<sup>^{10}</sup>$ To the best of our understanding of Pindyck's algorithm, estimates until time T and end-of-sample (up to time T) values of the state variables were used to compute out-of-sample forecasts.

<sup>&</sup>lt;sup>11</sup>See *e.g.* Schwartz and Smith (2000), Schwartz (1997), Cortazar and Schwartz (2003), and Bernard, Khalaf, Kichian and McMahon (2005).

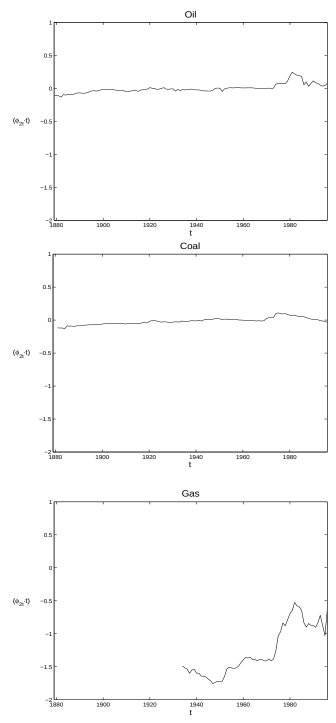


Figure 1. Estimated Price Trends

standard and depend on nuisance parameters. Using simulation-based procedures to address this issue, we have reported results for both a standard Monte Carlo test and a maximized Monte Carlo test. Our findings lend support to the proposed TVP class of energy models against the null hypothesis of fixed-coefficient mean-reverting equations. That is, we have found statistically significant instabilities for oil and natural gas prices, but not for coal prices. We have conducted out-of-sample forecasting exercises to differentiate between various mean-reverting and unit-root models. The TVP class emerges as the best fit for forecasting purposes.

Our main finding is that there is little evidence in favour of an increasing trend in the price of oil from 1890 to 1996, although OPEC decisions did leave an imprint in the late seventies. The same observations applies for the price of coal. However, the natural gas price series which started in 1919 displays a U-shaped trend which clearly slopes upward. Our results regarding the oil price trend agrees with the long held view of Adelman (2004): our estimated oil price trends shows no indication of impending scarcity but bears the marks of the OPEC decisions in the post 1973 era.

# Appendix

### A. Maximized Monte Carlo tests

Consider a (continuous) test statistic, S, whose distribution under the null hypothesis can be simulated once a finite set of nuisance parameters are specified. Conforming with the notational framework of Section 2, let us denote the nuisance parameter vector  $\theta \in \Omega$ , and let  $\Omega_0$  refer to the nuisance parameter subspace compatible with the null hypothesis  $H_0$  under test.

Denote by  $S_0$  the observed value of S and let  $S_j$ , j=1, ..., N refer to N i.i.d. random draws from the null distribution of the statistic (given the parameter value  $\theta$ ). A maximized Monte Carlo (MMC) test is defined by the critical region

$$\sup_{\theta \in \Omega_0} [\hat{p}_N(S_0 \mid \theta)] \le \alpha \tag{A.1}$$

where

$$\hat{p}_N(S_0 \mid \theta) = \frac{N\hat{G}_N(S_0 \mid \theta) + 1}{N+1}, \tag{A.2}$$

$$\hat{G}_{N}(S_{0} | \theta) = \frac{1}{N} \sum_{i=1}^{N} \mathbf{1}_{[0,\infty)}(S_{i} - S_{0}),$$

$$\mathbf{1}_{A}(x) = \begin{cases} 1, & \text{if } x \in A, \\ 0, & \text{if } x \notin A. \end{cases}$$
(A.3)

Note that  $N\hat{G}_N(S_0 \mid \theta)$  is the number of simulated values of the test statistic not smaller than  $S_0$ . The formula for  $\hat{p}_N(S_0 \mid \theta)$  gives a an *empirical p-value* for a given parameter vector  $\theta$ . Then we can show the MMC test based on (A.1) has level  $\alpha$  [see Dufour (2002)]:

$$\mathsf{P}\Big[\sup_{\theta \;\in\; \Omega_0} [\hat{p}_N(S_0 \,|\, \theta)] \leq \alpha\Big] \leq \alpha \quad \text{under $H_0$}.$$

The only condition needed to implement this procedure the possibility of simulating the relevant test statistic under the null hypothesis.  $^{12}$  The values of N and T (neither the number of replications nor the sample size) are taken as given, and no asymptotic argument is needed.

In this context, given any consistent estimate of  $\theta$  that satisfies  $H_0$  (denoted  $\hat{\theta}$ ), a parametric

<sup>&</sup>lt;sup>12</sup>For example, for  $\alpha=0.05$ , N can be as low as 19. Although, in principle, raising N will typically increase the test power and decrease its sensitivity to the underlying randomization, the simulation results reported in Dufour and Kiviet (1996, 1998), Dufour et al. (1998), Dufour and Khalaf (2001, 2002a, 2002b), and Dufour et al. (2004), suggest that increasing N beyond 99 has only a small effect on power.

bootstrap-type critical region can be obtained as:

$$\hat{p}_N(S_0 \mid \hat{\theta}) \le \alpha \,. \tag{A.4}$$

In general, however, nothing guarantees that the level property

$$P[\hat{p}_N(S_0|\theta) \leq \alpha] \leq \alpha \text{ under } H_0$$

holds. Under specific regularity conditions, the bootstrap p-value may be valid asymptotically in the sense that

$$\lim_{T \to \infty} \left\{ \mathsf{P}[\hat{p}_N(S_0|\hat{\theta}) \le \alpha] - \mathsf{P}[\hat{p}_N(S_0|\theta_0) \le \alpha] \right\} = 0 \tag{A.5}$$

where  $\hat{p}_N(S_0|\theta_0)$  is the empirical *p*-value that one would obtain for the "true" (unknown) nuisance parameters values. Generic conditions for (A.5) to hold are given in Dufour (2002).

Unfortunately, in the context of the TVP models, (A.5) may easily not hold, for example because of identification problems [see Dufour (1997, 2003)]. In practice, this means that, if a test based on (A.4) rejects, this result may be spurious *even in large samples*. Yet bootstrap non-rejections are not subject to the same limitations: if the bootstrap-type test is not significant, then we can be sure that the exact MMC test is not significant at level  $\alpha$ . Indeed,

$$\hat{p}_N(S_0|\hat{\theta}) > \alpha \Rightarrow \sup_{\theta \in \Omega_0} [\hat{p}_N(S_0|\theta)] > \alpha.$$

It is thus a good strategy to start the MMC sup-p-value step using a commonly used (e.g., a constrained QMLE) estimate of  $\theta$ .

## B. Kalman Filtering and time varying parameters

This appendix draws heavily on Kim and Nelson (1999, Chapter 3). Consider the TVP model (2.3), which we rewrite for convenience in matrix notation as:

$$y_t = H_t \beta_t + A z_t + \epsilon_t \,, \tag{B.1}$$

$$\beta_t = F\beta_{t-1} + \eta_t, \ t = 1, \dots, T,$$
 (B.2)

$$\epsilon_t \stackrel{i.i.d.}{\sim} N[0, R],$$
 (B.3)

$$\eta_t \stackrel{i.i.d.}{\sim} N[0, Q],$$
 (B.4)

where

$$y_t = P_t$$
,  $H_t = \begin{bmatrix} 1 & t \end{bmatrix}$ ,  $A = \begin{bmatrix} c_1 & c_2 \end{bmatrix}$ , (B.5)

$$\beta_t = \begin{bmatrix} \phi_{1t} \\ \phi_{2t} \end{bmatrix}, \quad z_t = \begin{bmatrix} 1 \\ P_{t-1} \end{bmatrix}, \quad z_t = \begin{bmatrix} 1 \\ P_{t-1} \end{bmatrix}, \quad \eta_t = \begin{bmatrix} v_{1t} \\ v_{2t} \end{bmatrix}, \quad (B.6)$$

$$F = \begin{bmatrix} c_3 & 0 \\ 0 & c_4 \end{bmatrix}, \quad Q = \begin{bmatrix} \sigma_{v_1}^2 & 0 \\ 0 & \sigma_{v_2}^2 \end{bmatrix}. \tag{B.7}$$

The prediction equations in the Kalman filter algorithm are given by:

$$\beta_{t|t-1} = F\beta_{t-1|t-1},$$
 (B.8)

$$\beta_{t|t-1} = F\beta_{t-1|t-1},$$

$$S_{t|t-1} = FS_{t-1|t-1}F' + Q,$$
(B.8)

where  $\beta_{t|t-1}$  is the forecast value of  $\beta_t$  on the basis of information available through date t-1, and  $S_{t|t-1}$  is its conditional variance. The conditional forecast error and its conditional variance can be obtained as:

$$e_{t|t-1} = y_t - H_t \beta_{t|t-1} - Az_t,$$
 (B.10)

$$f_{t|t-1} = H_t S_{t|t-1} H_t' + R.$$
 (B.11)

These expressions can be used in the updating equations of the algorithm according to

$$\beta_{t|t} = \beta_{t|t-1} + K_t e_{t|t-1},$$
 (B.12)

$$S_{t|t} = S_{t|t-1} - K_t H_t S_{t|t-1},$$
 (B.13)

where the Kalman gain term is  $K_t = S_{t-1|t-1} H_t' f_{t|t-1}^{-1}$ .

If, in addition to the error terms  $\epsilon_t$  and  $\eta_{it}$ , the initial value of  $\beta$  is Gaussian, then the distribution of  $y_t$  conditional on information available through time t-1 is also Gaussian, and its log-likelihood function is:

$$\ln(L) = -(1/2) \sum_{t=1}^{T} \ln(2\pi f_{t|t-1}) - (1/2) \sum_{t=1}^{T} e'_{t|t-1} f_{t|t-1} e_{t|t-1}.$$
 (B.14)

Therefore, given initial values for model parameters and state variables, the log-likelihood function can be maximized over the sample to yield maximum-likelihood parameter estimates. See Kim and Nelson (1999) for additional details.

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