Researchers employ the directional distance function (DDF) to estimate multiple-input and multiple-output production, firm inefficiency, and productivity growth. We relax restrictive assumptions by computing optimal directions subject to profit maximization and cost minimization, correct for the potential endogeneity of inputs and outputs, estimate latent prices for bad outputs, measure firms’ responses to shadow prices rather than actual prices, and introduce an unobserved productivity term into the DDF. For an unbalanced panel of U.S. electric utilities, a model assuming profit-maximization outperforms one assuming cost-minimization, while lagged productivity and energy price have the greatest effect on productivity.
1 Introduction

As developed by Caves et al. (1982a, 1982b), the distance function (DF) has been widely used to estimate radial representations of frontier production technologies where firms employ multiple good inputs to produce multiple good outputs. The distance from a production frontier is a measure of the firm’s technical efficiency (TE). The change in this measure over time is efficiency change (EC), while the shift in the frontier over time is technical change (TC). The sum of these two measures is productivity change (PC). The DF is input- (output-) oriented if all inputs (outputs) are proportionally scaled down (up) to reach the production frontier while all outputs (inputs) are held constant.

One major shortcoming of the DF is that an entire set of inputs or an entire set of outputs must be scaled by the same factor. This becomes problematic when modelling the generation of electricity, since good inputs (capital, labor, and energy) and bad inputs (such as sulfur) produce good outputs (residential and industrial/commercial electricity) and bad outputs (pollutants). Using the DF, the researcher is not able to differentially credit the firm for simultaneously reducing bad outputs while increasing good outputs. In response, many authors have estimated an output DF and treated bad outputs like good inputs (holding both constant). However, this does not credit the firm for reducing bad outputs. Also, if bad inputs are consumed, no credit is given for their reduction.\footnote{A bad input like sulfur would be consumed only when it is organically bound to the coal and oil which are burned to generate electricity. To our knowledge, only Yaisawarng and Klein (1994) include fuel sulfur content and sulfur dioxide emissions in a study of electric utility production.}

As an alternative, Chambers (1998) and Chambers et al. (1998) developed the directional distance function (DDF) which provides greater flexibility. It allows measurement of unique additive changes in each input and output through the calculation of different directions of movement for each to reach the production frontier. If non-zero directions are used to change only inputs (outputs), the DDF is input- (output-) oriented. When non-zero directions are used to change all inputs and outputs, the DDF is technology-oriented.

Despite the greater flexibility of the DDF, researchers typically impose three overly-restrictive assumptions. First, the researcher usually specifies arbitrary directions of movement of current firm production toward the frontier to measure inefficiency.\footnote{For example, assuming fixed directions, Färe et al. (2005) estimate an output DDF for electric utilities involving good inputs, a good output, and a bad output.} How-
ever, different directions of movement toward the frontier will generate different measures of inefficiency. Three Data Envelopment Analysis (DEA) studies seek to avoid arbitrary assignment of directions by using linear programming methods to choose directions that maximize the measured distance (i.e., technical inefficiency) of the firm relative to a DDF. The first, by Färe et al. (2013), considers only good inputs and good outputs. The second, by Hampf and Krüger (2015), extends this analysis by including bad outputs. The stated goal of the third paper, by Zofio et al. (2013), is to compute optimal directions consistent with a firm’s profit-maximization (PM) position on a DDF. They assume that firms are currently profit-maximizers and then measure the maximum distance from the current position. However, to measure the technology and productivity at the PM position, one must estimate the DDF jointly with the first-order conditions for PM. Since the latter are not included in their optimization model, the estimated directions cannot be consistent with PM. In this paper we estimate these conditions jointly with the DDF and compute directions consistent with PM, which we term “optimal-PM” directions.

Our approach follows Chambers (1998), who formulates a PM problem which includes a technology-oriented DDF (to measure the distance from the production frontier), and derives the first-order price equations for good inputs and outputs. In order to compute optimal-PM directions, Atkinson and Tsionas (2016) (AT) estimate the DDF jointly with the first-order price equations for only good inputs and good outputs, since the prices of bad outputs and bad inputs are missing. A complete set of utility-specific pollution permit prices (shadow prices for bad outputs) for the years of our sample data does not exist. As explained below, the prices of coal and oil include rebates for greater amounts of the bad input, sulfur. However, data is not publicly available to compute an hedonic price for sulfur.³ We generalize AT by assuming a data generating process for latent prices of regulated bad outputs. These latent prices replace missing actual prices, allowing us to add the first-order price equations for regulated bad outputs to the AT system.

The second restrictive assumption of many DDF models is that all input and output quantities are exogenous. Highly-influential papers by Olley and Pakes (1996) (OP) and Levinsohn and Petrin (2003) (LP) consider the problem of estimating productivity in the

³In the more typical industry study, prices of all inputs are missing and our methodology can be employed to estimate their first-order price equations having generated their estimated latent prices.
presence of endogenous inputs using panel data. Both papers estimate a single-output Cobb-Douglas production function with a two-component random error term. The first component is firm- and time-varying productivity that is unobserved by the econometrician but observed, at least in part, by the firm. Since the firm takes productivity into account to some degree in choosing its inputs, endogeneity results. The second random component is an idiosyncratic error that is assumed to be uncorrelated with the explanatory variables and the productivity component. With the OP approach, the econometrician proxies for the unobserved productivity component with a potentially observable function. To obtain this function, OP first specify that investment is a monotonic function of productivity for a given level of capital and vintage. They then invert this function to obtain the productivity component as a proxy function of capital, investment, and vintage. Following OP, LP replace investment with materials and solve for the productivity component as a proxy function of capital, materials, and vintage. Productivity is assumed to follow a first-order Markov process. After discussing the modification of OP and LP by Ackerberg, Caves, and Frazer (2015) (regarding when the firm chooses labor), Wooldridge (2009) provides the exact set of moment conditions required to identify each of these models, where instruments are subsets of current and lagged inputs. However, as Griliches and Mairesse (1998) stress, if the econometrician incorrectly specifies the productivity function, some degree of endogeneity remains. Both OP and LP recognize the possible invalidity of their instruments as well as the typical validity but unavailability of input and output prices as instruments.

In this paper, we avoid assuming that inputs are exogenous for electric utilities. In our sample, they vary input choice over time and these choices are arguably correlated with the idiosyncratic error term, when one misspecifies the proxy equation for productivity. This results in the endogeneity of input quantities. Such a result potentially applies to all input quantities with a cost-minimization (CM) model and to all input and output quantities with a PM model. Instead, we utilize the prices of good inputs and good outputs in our instrument set, since they are arguably exogenous. Utilities are price takers in input markets, since these markets are national (due to trans-continental oil and natural gas pipelines, trans-continental rail lines hauling coal and oil, and national mobility of labor and capital). Regulated utilities, which comprise the vast majority of our sample, face output prices that are set by regulatory commissions. The smaller number of restructured utilities face market-determined prices for good inputs and out-
puts. Thus, for both types of firms, we employ input and output prices rather than input quantities in our instrument set.

The third restrictive assumption with all previous DDF models is that actual prices equal shadow (perceived) prices for the firm. If the two sets of prices differ, the researcher must calculate optimal directions using shadow prices. Previous papers have developed the methodology to estimate shadow prices for profit, cost, and distance functions as summarized in Kumbhakar and Lovell (2000). However, our paper is the first to estimate shadow prices using a DDF and the first-order price equations from PM. We identify shadow prices by including input and firm-specific price inefficiency parameters in these equations. These parameters are estimated jointly with optimal-PM directions.

In addition, this paper is the first to estimate a model free of these three restrictive assumptions and, at the same time, explain the sources of firm productivity, without resorting to inconsistent two-step methods. Typically the two steps are: 1) regress output on a set of inputs and 2) regress the residuals on a set of explanatory variables that were omitted from the first step. The two sets of variables must be uncorrelated to avoid a potentially substantial bias. We avoid this improbable requirement by employing an unrestricted profit function from which we derive productivity as an estimable function of lagged productivity, profits, prices of inputs and outputs, vintage, and time. We include this measure of productivity as an input in the DDF. This enables us to compute the partial elasticities of productivity with respect to its arguments and decompose productivity growth.

We apply our methodology to an unbalanced panel of U.S. electric utilities. This sample significantly expands the AT data set by 80% to include years when a number of utilities were restructured. We report posterior densities for optimal directions, TE, EC, TC, PC, the resource implications of price inefficiency, and the sources of PC.

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4 The goal of deregulation was to increase competition, yielding greater TE, productivity growth, and price efficiency. On the production frontier, the profit-maximizing firm achieves price efficiency when the price of each input equals the value of its marginal product. The cost-minimizing firm achieves allocative efficiency when ratios of input prices equal ratios of their marginal products.

5 Reasons for deviations of shadow from actual prices include tax write-offs, rate-of-return regulation, and constraints imposed by regulatory agencies or labor unions.

6 See Wang and Schmidt (2002) for details on Monte Carlo experiments indicating substantial potential bias in both steps.
2 The Directional Distance Function

2.1 Computing Optimal Directions

We assume a firm production technology that combines good inputs, \( x = (x_1, \ldots, x_N) \in R^N_+ \), and bad inputs, \( \tilde{x} = (\tilde{x}_1, \ldots, \tilde{x}_J) \in R^J_+ \), to produce good outputs, \( y = (y_1, \ldots, y_M) \in R^M_+ \), and bad outputs, \( \tilde{y} = (\tilde{y}_1, \ldots, \tilde{y}_L) \in R^L_+ \). A firm with vintage, \( \tau \), productivity shock, \( \omega \), at time \( t = 1, \ldots, T \), has production technology

\[
T(\omega, \tau, t) = \{ (x, \tilde{x}, y, \tilde{y}; \omega, \tau, t) : x, \tilde{x} \text{ can produce } y, \tilde{y} \text{ with } (\omega, \tau, t) \}
\]  

(1)

Let \( g = (g_x, g_{\tilde{x}}, g_y, g_{\tilde{y}}) \) be a direction vector. Typically researchers assume that \( (g_x, g_{\tilde{x}}, g_y, g_{\tilde{y}}) = (-1, -1, 1, -1) \). Following Chambers (1998), we define the technology DDF as

\[
\overrightarrow{D}_T(x, \tilde{x}, y, \tilde{y}; \omega, \tau, t) = \sup \{ \beta : (x + \beta g_x, \tilde{x} + \beta g_{\tilde{x}}, y + \beta g_y, \tilde{y} + \beta g_{\tilde{y}}) \in T(\omega, \tau, t) \},
\]  

(2)

That is, the typical assumption is that the analyst measures the distance from the frontier using equal absolute values for directions which increase good outputs and reduce all other inputs and outputs.

We now specify the important properties of the technology DDF that hold whether directions are assigned or estimated optimally. As shown in Hudgins and Primont (2007), for any values of the elements of \( g \):

D1. Translation Property:

\[
\overrightarrow{D}_T(x + \alpha g_x, \tilde{x} + \alpha g_{\tilde{x}}, y + \alpha g_y, \tilde{y} + \alpha g_{\tilde{y}}; \omega, \tau, t) = \overrightarrow{D}_T(x, \tilde{x}, y, \tilde{y}; \omega, \tau, t) - \alpha,
\]  

(3)

D2. \( g \)-Homogeneity of Degree Minus One:

\[
\overrightarrow{D}_T(x, \tilde{x}, y, \tilde{y}; \lambda g_x, \lambda g_{\tilde{x}}, \lambda g_y, \lambda g_{\tilde{y}}; \omega, \tau, t) = \lambda^{-1} \overrightarrow{D}_T(x, \tilde{x}, y, \tilde{y}; g, \omega, \tau, t), \quad \lambda > 0,
\]  

(4)

D3. Concavity:

\[
\overrightarrow{D}_T(x, \tilde{x}, y, \tilde{y}; g, \omega, \tau, t) \text{ is concave in } (x, \tilde{x}, y, \tilde{y}; g, \omega, \tau, t),
\]  

(5)

D4. Non-negativity:

\[
\overrightarrow{D}_T(x, \tilde{x}, y, \tilde{y}; g, \omega, \tau, t) \geq 0, \quad (x, \tilde{x}, y, \tilde{y}; g, \omega, \tau, t) \in T(\omega, \tau, t).
\]  

(6)

Equation (3) says that the technology DDF will satisfy the translation property. For example, increasing \( y \) and decreasing \( x, \tilde{x}, \) and \( \tilde{y} \) by \( \alpha \), each multiplied by their direction,
will result in a decrease in the technology DDF by $\alpha$. This is analogous to the property of linear homogeneity with a Shephard distance function. Equation (4) indicates that scaling each direction by $\lambda$ will scale the technology DDF by $\lambda^{-1}$. Equation (5) imposes concavity of the technology DDF. Finally, equation (6) requires that the technology DDF function be non-negative, which is easily imposed after estimation.

For the following properties we let “S” represent the assumption of strong disposability of all inputs and outputs:

**D5–S. Good Input Monotonicity:** We first assume that good inputs are strongly disposable. In Appendix A.1 we show that this implies

$$\frac{\partial \tilde{D}_T(x, \tilde{x}, y, \tilde{y}; g, \omega, \tau, t)}{\partial x_n} \geq 0, n = 1, \ldots, N. \tag{7}$$

**D6–S. Good Output Monotonicity:** We assume that good outputs are strongly disposable. Following the proof of D5–S, in Appendix A.2, we show that this assumption implies

$$\frac{\partial \tilde{D}_T(x, \tilde{x}, y, \tilde{y}; g, \omega, \tau, t)}{\partial y_m} \leq 0, m = 1, \ldots, M. \tag{8}$$

Following the proof of D5–S, assuming that bad inputs are strongly disposable, we can determine

**D7–S. Bad Input Monotonicity:**

$$\frac{\partial \tilde{D}_T(x, \tilde{x}, y, \tilde{y}; g, \omega, \tau, t)}{\partial \tilde{x}_j} \geq 0, j = 1, \ldots, J. \tag{9}$$

Again following the proof of D5–S, assuming that bad outputs are strongly disposable, we can determine

**D8–S. Bad Output Monotonicity:**

$$\frac{\partial \tilde{D}_T(x, \tilde{x}, y, \tilde{y}; g, \omega, \tau, t)}{\partial \tilde{y}_l} \geq 0, l = 1, \ldots, L. \tag{10}$$

However, researchers typically do not assume that bads are strongly disposable. Instead, most investigators maintain that bad inputs and bad outputs are weakly disposable with good inputs and outputs. The weak disposability of bad inputs results from their organic combination with good inputs, so that a reduction in bad inputs implies a reduction in good inputs, outputs held constant. Likewise, bad outputs are weakly disposable since to reduce them, we must divert resources from the production of good outputs, holding inputs constant. Assuming weak disposability, we cannot determine the global monotonicity property of bad inputs and bad outputs as in D7–S and D8–

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5This paper with all Appendices is stored at RePec.
S. However, as shown shortly, if we assume weak disposability, we can determine local monotonicity conditions given that we also assume PM and know the signs of the prices of bad inputs and bad outputs. If instead we assume CM, the firm is subject to only the first-order conditions for inputs, so that the monotonicity conditions for good and bad inputs are the same as with the PM model.

A number of studies have modeled electric utilities assuming PM. See Atkinson and Halvorsen (1976, 1980) and Cowing (1978). Other researchers have assumed CM subject to a set of regulatory constraints on earned rates of return and a requirement to satisfy all demand at a given price (that is, output is taken as given). However, output may be endogenous as with an ex ante cost function. Further, if these regulatory constraints are not binding, utilities may maximize profits. Fowlie (2010) provides evidence of this by showing that many regulated utilities earn allowed rates of return on capital that considerably exceed the market rate of return, indicating that constraints on profits may not be binding and output may be endogenous. These results indicate that a PM model may be more appropriate than a CM model. Thus, we focus on the PM model and compare the accuracy of its results to those of the CM model.

Temporarily suppressing all the arguments of \( \overrightarrow{D_T}(x, \tilde{x}, y, \tilde{y}; g, \omega, \tau, t) \), we follow Chambers (1998) and assume that a firm maximizes profits, \( \pi \), by choosing values of \( x, y, \tilde{x}, \tilde{y} \) to solve

\[
\sup \left\{ p_y (y + \overrightarrow{D_T}g_y) - p_y (\tilde{y} + \overrightarrow{D_T}g_{\tilde{y}}) - p_x (x + \overrightarrow{D_T}g_x) - p_{\tilde{x}} (\tilde{x} + \overrightarrow{D_T}g_{\tilde{x}}) \right\}
\]

where \( p_y \geq 0, p_{\tilde{y}} \geq 0, p_x \geq 0, \) and \( p_{\tilde{x}} \leq 0 \) are price vectors and the econometrician either pre-determines or estimates \( g \). Further, we define \( p = (p_y, p_{\tilde{y}}, p_x, p_{\tilde{x}}) \).

Typically (as indicated above) the DDF is estimated without the first-order conditions for PM, where one has assumed a set of \textit{a priori} fixed directions. However, in this paper we assume that the firm chooses \( (x, \tilde{x}, y, \tilde{y}) \) subject to the first-order conditions for PM and we estimate optimal-PM directions consistent with these conditions. The

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\( ^8 \)See Pope and Just (1996) regarding identification and estimation of an ex ante cost function.

\( ^9 \)The CM model is obtained by using only the first-order conditions for input prices and assuming that output is given.
first-order conditions are:

\[ p_n / \varrho(p, g) = \partial \overrightarrow{D}(x, \tilde{x}, y, \tilde{y}; g, \omega, \tau, t) / \partial x_n, n = 1, \ldots, N, \]  

\[ p_m / \varrho(p, g) = -\partial \overrightarrow{D}(x, \tilde{x}, y, \tilde{y}; g, \omega, \tau, t) / \partial y_m, m = 1, \ldots, M, \]  

\[ p_j / \varrho(p, g) = \partial \overrightarrow{D}(x, \tilde{x}, y, \tilde{y}; g, \omega, \tau, t) / \partial \tilde{x}_j, j = 1, \ldots, J, \]  

\[ p_l / \varrho(p, g) = \partial \overrightarrow{D}(x, \tilde{x}, y, \tilde{y}; g, \omega, \tau, t) / \partial \tilde{y}_l, l = 1, \ldots, L, \]

where

\[ \varrho(p, g) = [\sum m p_m g_m - \sum n p_n g_n - \sum j p_j g_j - \sum l p_l g_l]. \]

We assume that \( \varrho(p, g) > 0 \), where \( \varrho(p, g) \) is the optimal value of the Lagrangian multiplier, which is the change in profits due to a small improvement in the production technology. For details see Hudgins and Primont (2007) who show that one can solve the profit-maximization problem in (11) or solve the equivalent Lagrangian function

\[ L = p_y y - p_x x - p_{\tilde{y}} \tilde{y} - p_{\tilde{x}} \tilde{x} + \varrho \overrightarrow{D}(x, \tilde{x}, y, \tilde{y}; g, \omega, \tau, t), \]

in order to obtain this interpretation of \( \varrho \).

All prices are assumed to be non-negative except for the price of bad inputs, which is non-positive. The prices of good outputs and the prices of good inputs are non-negative by definition. The price of bad outputs is positive, since the firm must pay a fine or buy emission permits for additional production of bad outputs. When a bad input is organically bound to a good input, the price of the bad input is negative, since the firm must be compensated for utilizing it.

For the following two properties, “W” indicates weakly disposable. Assuming weak disposability of bad inputs and that \( p_{\tilde{x}} \leq 0 \), from (14) we obtain locally:

**D7–W.** Bad Input Monotonicity:

\[ \partial \overrightarrow{D}(x, \tilde{x}, y, \tilde{y}; g, \omega, \tau, t) / \partial \tilde{x}_j \leq 0, j = 1, \ldots, J. \]

Assuming weak disposability of bad outputs and that \( p_{\tilde{y}} \geq 0 \) from (15) we obtain locally:

**D8–W.** Bad Output Monotonicity:

\[ \partial \overrightarrow{D}(x, \tilde{x}, y, \tilde{y}; g, \omega, \tau, t) / \partial \tilde{y}_l \geq 0, l = 1, \ldots, L. \]

In this manner we maintain PM and use assumptions about the prices of bad inputs and bad outputs to locally restrict the range of the partial derivatives in D7–W and D8–W. For the CM model, only the monotonicity conditions for good and bad inputs apply. They are the same as with the PM model.
Rather than assume fixed directions, in this paper we compute optimal directions that are consistent with PM and CM. In Fig. 1, a firm increases a good output and reduces a bad output by moving from $z$ to $z^*$, the PM point. This is consistent with the traditional assumptions, where the directions for the good output and bad output are positive and negative, respectively. However, the signs of optimal directions may be quite different. In Fig. 1, if a firm moved from $z$ to the point of PM at $z'$, from (2) the directions $(g_y, g_{\tilde{y}})$ are both positive since the good output and the bad output would both increase. One can easily substitute $\tilde{x}$ or $x$ for $\tilde{y}$ and obtain similar results.

An electric utility subject to the first-order conditions for PM may maximize profits by increasing good outputs, increasing some inputs to produce the additional good outputs, decreasing regulated bad outputs, and increasing unregulated bad outputs due to the increase in good outputs. Thus, we impose no a priori sign restrictions on the optimal direction of any input or output.

The input and output DDF are easily obtained as special cases of the technology DDF. The output DDF changes good and bad outputs in the direction $(0, 0, g_y, g_{\tilde{y}})$, for a given level of inputs in order to move to the frontier of $T(\omega, \tau, t)$. The input DDF changes good and bad inputs in the direction $(g_x, g_{\tilde{x}}, 0, 0)$, for a given level of good and bad outputs in order to move to the frontier of $T(\omega, \tau, t)$.

3 Econometric Formulation

3.1 The Technology Directional Distance System and the Translation Restrictions

Assume that we have panel data for firm $i (i = 1, \ldots, N)$ in time period $t (t = 1, \ldots, T)$ on all inputs and outputs. We then formulate our technology DDF as a quadratic function of $x, \tilde{x}, y, \tilde{y}, \tau,$ and $t$ as:

$$
\overline{D}_T(x_{it}, \tilde{x}_{it}, y_{it}, \tilde{y}_{it}; \tau_{it}, t) = \sum_{n=1}^{N} \gamma_n(x_{n,it}) + \sum_{m=1}^{M} \gamma_m(y_{m,it}) + \sum_{j=1}^{J} \gamma_j(\tilde{x}_{j,it}) + \sum_{l=1}^{L} \gamma_l(\tilde{y}_{l,it})
+ \frac{1}{2} \sum_{n=1}^{N} \sum_{n'=1}^{N} \gamma_{nn'}(x_{n,it})(x_{n',it}) + \frac{1}{2} \sum_{j=1}^{J} \sum_{j'=1}^{J} \gamma_{jj'}(\tilde{x}_{j,it})(\tilde{x}_{j',it})
+ \frac{1}{2} \sum_{m=1}^{M} \sum_{m'=1}^{M} \gamma_{mm'}(y_{m,it})(y_{m',it}) + \frac{1}{2} \sum_{l=1}^{L} \sum_{l'=1}^{L} \gamma_{ll'}(\tilde{y}_{l,it})(\tilde{y}_{l',it})
$$
\[ \sum_{j=1}^{J} \sum_{n=1}^{N} \gamma_{jn}(\tilde{x}_{j,it})(x_{n,it}) + \sum_{m=1}^{M} \sum_{n=1}^{N} \gamma_{mn}(y_{m,it})(x_{n,it}) \\
+ \sum_{l=1}^{L} \sum_{n=1}^{N} \gamma_{ln}(\tilde{y}_{l,it})(x_{n,it}) + \sum_{m=1}^{M} \sum_{j=1}^{J} \gamma_{jm}(\tilde{x}_{j,it})(y_{m,it}) \\
+ \sum_{l=1}^{L} \sum_{j=1}^{J} \gamma_{jl}(\tilde{x}_{j,it})(y_{m,it}) + \sum_{m=1}^{M} \sum_{l=1}^{L} \gamma_{lm}(\tilde{y}_{l,it})(y_{m,it}) \\
+ \sum_{n=1}^{N} \gamma_{nt}(x_{n,it})d_t + \sum_{m=1}^{M} \gamma_{mt}(y_{m,it})d_t + \sum_{j=1}^{J} \gamma_{jt}(\tilde{x}_{j,it})d_t \\
+ \sum_{l=1}^{L} \gamma_{lt}(\tilde{y}_{l,it})d_t + \sum_{t=1}^{T} \gamma_{t}(x_{n,it})\tau_{it} \\
+ \sum_{m=1}^{M} \gamma_{mr}(y_{m,it})\tau_{it} + \sum_{j=1}^{J} \gamma_{jr}(\tilde{x}_{j,it})\tau_{it} \\
+ \sum_{l=1}^{L} \gamma_{lr}(\tilde{y}_{l,it})\tau_{it} + \gamma_{r}\tau_{it}, \tag{20} \]

where \( d_t \) is a year dummy. We later accept the null that \( d_t \) and \( \tau_{it} \) enter (20) linearly using a likelihood ratio test at the .05 level. Hence, we drop all interaction terms involving these variables from this equation.\(^{10}\)

We can now restate the first-order conditions for PM, (12)-(15), in terms of the parameters of the quadratic DDF for each good input price equation as

\[ \frac{p_{n,it}}{\psi} = \gamma_n + \sum_{n'=1}^{N} \gamma_{nn'}(x_{n',it}) + \sum_{j=1}^{J} \gamma_{jn}(\tilde{x}_{j,it}) + \sum_{m=1}^{M} \gamma_{mn}(y_{m,it}) + \sum_{l=1}^{L} \gamma_{ln}(\tilde{y}_{l,it}), \tag{21} \]

for each good output price equation as

\[ \frac{p_{m,it}}{\psi} = -\left[ \gamma_m + \sum_{m'=1}^{M} \gamma_{mm'}(y_{m',it}) + \sum_{j=1}^{J} \gamma_{jm}(\tilde{x}_{j,it}) + \sum_{n=1}^{N} \gamma_{mn}(x_{n,it}) + \sum_{l=1}^{L} \gamma_{lm}(\tilde{y}_{l,it}) \right], \tag{22} \]

for each bad input price equation as

\[ \frac{p_{j,it}}{\psi} = \gamma_j + \sum_{j'=1}^{J} \gamma_{jj'}(\tilde{x}_{j',it}) + \sum_{n=1}^{N} \gamma_{jn}(x_{n,it}) + \sum_{m=1}^{M} \gamma_{jm}(y_{m,it}) \]

\(^{10}\)Färe and Lundberg (2005) prove that only two functional forms have a second-order Taylor series approximation interpretation of the DDF and satisfy the translation property. These are the logarithmic transcendental and the quadratic. Also see Chambers (1998) for further discussion of this point. We employ the quadratic since it is linear in the parameters.
and for each bad output price equation as

\[ p_{l,it}/\varrho = \gamma_l + \sum_{l'=1}^{L} \gamma_{l'l} (\tilde{y}_{l',it}) + \sum_{j=1}^{J} \gamma_{jl} (\tilde{x}_{j,it}) + \sum_{n=1}^{N} \gamma_{ln} (x_{n,it}) + \sum_{m=1}^{M} \gamma_{lm} (y_{m,it}). \]  

(24)

The restrictions guaranteeing the translation property in (3) are imposed parametrically on (20) and on (21)-(24). To derive these restrictions for an input, an output, and a technology DDF, we generalize Hudgins and Primont (2007) by adding bad inputs and bad outputs. To simplify notation, let \( \tilde{z} = (x, \tilde{x}, y, \tilde{y}) \). First, assuming a quadratic functional form for the technology DDF:

\[ \tilde{D}_T(\tilde{z}) = \sum_{w=1}^{W} \gamma_w \tilde{z}_w + \sum_{w=1}^{W} \sum_{w'=1}^{W} \gamma_{ww'} \tilde{z}_w \tilde{z}_{w'} , \]  

(25)

where \( w = 1, \ldots, W, W = M + N + J + L \). To determine the appropriate parametric restrictions that guarantee the translation property incorporating \( g \), we note that the translation property requires that

\[ \tilde{D}_T(\tilde{z} + \alpha g) = \sum_{w=1}^{W} \gamma_w (\tilde{z}_w + \alpha g_w) + \sum_{w=1}^{W} \sum_{w'=1}^{W} \gamma_{ww'} (\tilde{z}_w + \alpha g_w) (\tilde{z}_{w'} + \alpha g_{w'}) = \sum_{w=1}^{W} \gamma_w \tilde{z}_w + \sum_{w=1}^{W} \sum_{w'=1}^{W} \gamma_{ww'} \tilde{z}_w \tilde{z}_{w'} - \alpha. \]  

(26)

### 3.2 The Stochastic Framework with Shadow and Latent Prices

#### 3.2.1 Stochastically Imposing the Translation Property Restrictions

Generalizing Hudgins and Primont (2007), we derive the following parametric restrictions to stochastically impose the translation property in (26) for the technology

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11 An alternative to the quadratic is the logarithmic transcendental technology translation function which automatically satisfies the translation properties. See Chambers (1998) for more details. Note that we can approximate a DDF with a quadratic function, but not a translog. Using the former, one can impose the translation property, since original variables are used. However, one cannot impose the translation property on the latter.
where $\vartheta_o \sim \mathcal{N}(-1, c^2)$ and $\vartheta \sim \mathcal{N}(0, c^2)$, $\mathcal{N}$ is the normal density, and $c = 10^{-4}$ to keep the variance small around zero. The stochastic restrictions in (27) are, in fact, semi-informative priors placed upon $g$ and $\gamma$, a vector of all $\gamma_w$ parameters. These parameters are estimated jointly, as explained below. In a more traditional, non-Bayesian approach, one would set $\vartheta_o = -1, \vartheta = 0$.

To obtain the corresponding restrictions for the input and output DDF models, simply eliminate the summation terms for the outputs and the inputs, respectively. For the technology DDF model, we have imposed symmetry for all the double-subscripted coefficients for all inputs and outputs. Note that the introduction of parameters to measure the direction for each input and output occurs only through the translation property restrictions.

### 3.2.2 Modeling $\omega$

Assuming now that production is a function of inputs, outputs, vintage and a Hicks-neutral productivity shock, $\omega$, the resulting profit function obtained from (11) is

$$\pi = \pi(p_y, p_{\tilde{y}}, p_x, p_{\tilde{x}}, \omega; g, \tau, t) = \pi(p, \omega; g, \tau, t).$$  \hspace{1cm} (28)

By definition, $\pi$ is non-decreasing, monotonic in $\omega$. Strengthening this condition to positive monotonicity of $\pi$ in $\omega$, we can write the inverse function for $\omega$ as $\omega = f(p, \pi; g, \tau, t)$.

Subject to (27) we specify a stochastic framework for (20) as:
\[ 0 = \overrightarrow{D}_T(x, \tilde{x}, y, \tilde{y}; g, \tau, t) + v_{it} + \omega_{it} - u_{it}, \]  
(29)

where the stochastic part is comprised of an idiosyncratic i.i.d. term, \( v_{it} \), which has zero mean, \( \omega_{it} \), and a one-sided component, \( u_{it} \). While \( v_{it} \) reflects errors in optimization due to random events beyond the control of the firm (such as weather), \( u_{it} \) reflects firm-specific inefficiencies that may vary over time. We generalize the productivity component, \( \omega_{it} \), by including \( \omega_{i,t-1} \), which is lagged \( \omega_{it} \), and utilizing \( p^*_i \), where the star indicates that latent prices replace missing prices:

\[ \omega_{it} = f(\omega_{i,t-1}, p^*_i, \pi_{it}; g, \tau_{it}, t) + \epsilon_{it}, \]  
(30)

and we must obtain an approximation to the unknown functional form \( f(\cdot) \). We further specify that

\[ \log(u_{it}) = \gamma_1 + \gamma_2 \omega_{it} + \gamma_3 \omega_{i,t-1} + \gamma_4 \log u_{i,t-1} + \gamma_5 t + d^\prime_{it} + Z^\prime_{i,t-1} \delta + \epsilon_{it,2}, \]  
(31)

where \( d_{it} \) denotes firm dummies, and \( Z_{i,t-1} \) contains lagged values of all inputs and outputs.

We now consider different approximations of (30). To obtain a translog-neural-network approximation, let \( z_{it} = (\omega_{i,t-1}, p^*_i, \pi_{it}; g, \tau_{it}, t) \). Then

\[ f(z_{it}) = a_0 + a^\prime z_{it} + \frac{1}{2} z_{it}^\prime A z_{it} + \sum_{g=1}^{G} \lambda_g \varphi(z_{it}^\prime b_g), \]  
(32)

where the activation function \( \varphi(\kappa) = \frac{1}{1+\exp(-\kappa)}, -\infty < \kappa < \infty \). As a second alternative, we use a second-order approximation:

\[ f(z_{it}) = a_0 + a^\prime z_{it} + \frac{1}{2} z_{it}^\prime A z_{it}. \]  
(33)

As a third alternative, we use the Fourier approximation:

\[ f(z_{it}) = a_0 + a^\prime z_{it} + \frac{1}{2} z_{it}^\prime A z_{it} + \sum_{\phi=1}^{\Phi} \left\{ u_{\Phi 0} + 2 \sum_{\ell=1}^{J} \left( u_{\ell \Phi} \cos(k_{\ell \Phi}^\prime z_{it}) + v_{\ell \Phi} \sin(k_{\ell \Phi}^\prime z_{it}) \right) \right\}, \]  
(34)

where \( k_\phi \) is a multi-index, \( \Phi \) is a number determined by \( \text{dim}(z_{it}) \), \( J \) is the order of the expansion, and \( u_{\Phi 0}, u_{\ell \Phi}, v_{\ell \Phi} \) are unknown parameters. The multi-indices are constructed using the following rules: i) the zero vector and any \( k_\phi \) whose first non-zero element is negative are deleted; ii) Every index with a common integer divisor is also deleted. Gallant (1982) shows that \( A = -\sum_{\phi=1}^{\Phi} u_{\Phi 0} k_\phi k_\phi^\prime \). Notice that we can leave \( A \) unrestricted and obtain a different Fourier approximation. We use the same re-scaling as in Gallant (1982) and Feng and Serletis (2009) with the exception that we do not use logs. Finally, the purpose of the re-scaling is that elements of \( z_{it} \) must lie in \([0, 2\pi]\) which can
be achieved through a common scale in \( z_{it} \). Finally, we can employ a full third-order approximation:

\[
    f(z_{it}) = a_0 + \sum_{i=1}^{b} a_i z_{it} + \sum_{i=1}^{b} \sum_{j=1, i \leq j}^{b} b_{ij} z_{it} z_{jt} + \sum_{i=1}^{b} \sum_{j=1}^{b} \sum_{k=1, i \leq j \leq k}^{b} c_{ijk} z_{it} z_{jt} z_{kt},
\]

where \( b \) is the number of elements in \( z_{it} \).

We perform posterior analysis with all these specifications, which are flexible enough to cover nearly every empirical case. For any model with parameters \( \theta \in \Theta \subseteq \mathbb{R}^d \), including any latent variables in the model, denote the prior by \( p(\theta) \), the likelihood by \( \mathcal{L}(\theta; Y) \) for data \( Y \) and the posterior by \( p(\theta|Y) \). We know

\[
    p(\theta|Y) = \frac{\mathcal{L}(\theta; Y)p(\theta)}{p(Y)},
\]

where the marginal likelihood is

\[
    p(Y) = \int_\Theta \mathcal{L}(\theta; Y)p(\theta)d\theta.
\]

For two different models, say 1 and 2, we define the Bayes factor in favor of model 1 and against model 2 as:

\[
    BF_{1:2} = \frac{p_1(Y)}{p_2(Y)}.
\]

The Bayes factors, in our application, are reported in Table 2. These results clearly favor the third-order approximation in (35). All of our results will, therefore, be conditional on the selection of this functional form for the productivity equation.

The computation of the marginal likelihood is an involved operation. Here, we compute it using the importance sampling techniques in Perrakis, Ntzoufras and Tsionas (2014).

We also treat firm-specific directions, \( g_i \), as latent variables and assume the following prior:

\[
    g_i \sim \mathcal{N}_{N+M+J+L}(\bar{g}, \Sigma_g),
\]

where \( \bar{g} \) denotes the prior mean vector whose elements consist of reasonable prior beliefs (namely, -1 for all inputs and the bad outputs and +1 for the good outputs) and \( \Sigma_g \) is constructed so as to reflect reasonable deviations from these prior beliefs. Below we find that the posteriors are highly insensitive to variations in these priors.

Having estimated productivity (\( \omega_{it} \)), we compute \( PC = \frac{\partial \omega_{it}}{\partial t} \) and \( EC = \frac{\partial u_{it}}{\partial t} \), using equations (30) and (31). Then \( TC = PC - EC \). For more details on computing these quantities, see Grosskopf (2003). We also compute returns to scale (RTS) as the sum of the good output elasticities calculated from the distance function.
3.2.3 Shadow Prices

To incorporate shadow prices as the relevant prices for the firm we write:

\[
p^s_n = \frac{\partial \tilde{D}_T(z; g)}{\partial x_n}, \quad n = 1, \ldots, N, \quad \text{(40)}
\]

\[
p^s_m = -\frac{\partial \tilde{D}_T(z; g)}{\partial y_m}, \quad m = 1, \ldots, M, \quad \text{(41)}
\]

\[
p^s_j = \frac{\partial \tilde{D}_T(z; g)}{\partial \tilde{x}_j}, \quad j = 1, \ldots, J, \quad \text{(42)}
\]

\[
p^s_l = \frac{\partial \tilde{D}_T(z; g)}{\partial \tilde{y}_l}, \quad l = 1, \ldots, L. \quad \text{(43)}
\]

with \( g^s = \sum_{m=1}^M p^s_m g_m - \sum_{n=1}^N p^s_n g_n - \sum_{j=1}^J p^s_j g_j - \sum_{l=1}^L p^s_l g_l \), where the superscript \( s \) denotes shadow (or perceived) prices to the firm. The shadow prices satisfy the first-order conditions, which are used to complete the system, since we have many endogenous variables but only one DDF equation. Shadow prices are the relevant prices for the shadow-profit-maximizing firm. Actual prices potentially differ from shadow prices by an amount \( \xi_q, q = n, m, j, l \):

\[
p^s_n = p_n + \xi_n, \quad n = 1, \ldots, N, \quad \text{(44)}
\]

\[
p^s_m = p_m + \xi_m, \quad m = 1, \ldots, M, \quad \text{(45)}
\]

\[
p^s_j = p_j + \xi_j, \quad j = 1, \ldots, J, \quad \text{(46)}
\]

\[
p^s_l = p_l + \xi_l, \quad l = 1, \ldots, L. \quad \text{(47)}
\]

The variable \( \xi_{it} \) is latent, where we assume the following prior:

\[
\xi_{it} = [\xi_N, \xi_M', \xi_J', \xi_L']' \sim N_{N+M+J+L}(0, \Omega), \quad \text{(48)}
\]

where \( \xi_N = [\xi_{n,it}, n = 1, \ldots, N]' \), \( \xi_M = [\xi_{m,it}, m = 1, \ldots, M]' \), \( \xi_J = [\xi_{j,it}, j = 1, \ldots, J]' \), \( \xi_L = [\xi_{l,it}, l = 1, \ldots, L]' \), and where \( \Omega \) is unknown.

In view of (40)–(43), equations (44)–(47) are rewritten, after introducing stochastic...
error terms  \( v_q, q = n, m, j, l \), as:

\[
p_n = \varphi^s(p, g, \xi) \cdot \frac{\partial \hat{D}_T(z; g)}{\partial x_n} - \xi_n + v_n, n = 1, \ldots, N, \tag{49}
\]

\[
p_m = -\varphi^s(p, g, \xi) \cdot \frac{\partial \hat{D}_T(z; g)}{\partial y_m} - \xi_m + v_m, m = 1, \ldots, M, \tag{50}
\]

\[
p_j = \varphi^s(p, g, \xi) \cdot \frac{\partial \hat{D}_T(z; g)}{\partial \tilde{x}_j} - \xi_j + v_j, j = 1, \ldots, J, \tag{51}
\]

\[
p_l = \varphi^s(p, g, \xi) \cdot \frac{\partial \hat{D}_T(z; g)}{\partial \tilde{y}_l} - \xi_l + v_l, l = 1, \ldots, L, \tag{52}
\]

where \( \varphi^s(p, g, \xi) = \left( \varphi(p, g) + \sum_{m=1}^{M} \xi_m g_m - \sum_{n=1}^{N} \xi_n g_n - \sum_{j=1}^{J} \xi_j g_j - \sum_{l=1}^{L} \xi_l g_l \right) \), the vector \( \xi = (\xi_N, \xi_M, \xi_J, \xi_L) \), and the vector \( p = (p_N, p_M, p_J, p_L) \). We can normalize the first element of \( \xi_N \) to zero, since we can only identify relative price distortions.\(^{12}\)

Assuming that we have data on prices for all inputs and outputs, estimating all of the normalized price equations in (12)–(15) together with the DDF would generate a singularity in the covariance matrix of the residuals. This occurs since the normalized price equations sum to 1 for any value of \( g \) (as is seen from adding (12)-(15) with appropriate changes in sign and comparing to (16) ). The choice of which price equation to drop does not impact the results. The technology system is the DDF in (20) substituted into (29) and \( N + M + J + L - 1 \) of the price equations in (49)–(52), subject to imposition of the restrictions in (27). Relative to one normalized input or output efficiency parameter, we can estimate both input and output price efficiencies by estimating this system, since all the price equations are included.

### 3.2.4 Completing the System using Reduced-Form Equations

Assume now a worse-case scenario where we lack prices and alternative valid instruments for the endogenous bads, \( \tilde{x} \) and \( \tilde{y} \), so that we are unable to identify the DDF using equations like (51) and (52). Assume further that we are unable to generate latent prices for \( \tilde{x} \) and \( \tilde{y} \). We can still identify the technology DDF. First we create a system of equations consisting of (20) substituted into (29) and \( N + M + J + L - 1 \) of the price equations in (49) and (50), subject to the restrictions in (27). Then we complete this system by including reduced-form equations for \( \tilde{x} \) and \( \tilde{y} \) following standard LIML practices as in

\(^{12}\)This occurs since PM means that the firm chooses absolute levels of outputs given CM, which requires that the firm equate ratios of input prices to ratios of marginal products. Estimation of the extent to which the latter has been achieved requires one normalization.
Pagan (1979). The input and output DDF systems would be constructed analogously. The explanatory variables for these equations are prices of the good inputs and good outputs, firm dummies, and time dummies. See Appendix B for further details.

### 3.2.5 Completing the System using Latent-Price Equations

Frequently, actual prices for inputs and outputs are missing. With rare exceptions, prices for good inputs are confidential for privately-owned, unregulated firms. While data on prices and quantities of good inputs and outputs are typically reported by regulated electric utilities, the federal government currently allows many utilities to redact data on wages. Publicly-available data sources intermittently report prices from thin tradable permit markets for the regulated bad outputs that we examine, sulfur dioxide (SO$_2$) and nitrogen oxide (NO$_x$). No reliable prices exist for the unregulated pollutant that we model, carbon dioxide (CO$_2$). Further, prices of bad inputs are missing for all firms. In theory one could estimate prices for sulfur, a bad input, which is chemically bound to coal and oil, which are good fuel inputs. However, this would require the use of hedonic methods, which are infeasible due to data confidentiality.\(^\text{13}\)

In place of missing actual prices, we can sample prices from their conditional posteriors generated by their FOCs to generate latent prices $p^*_j$ and $p^*_l$ for $\tilde{x}$ and $\tilde{y}$ to replace the missing $p_j$ and $p_l$ in (51) and (52), respectively. Then we can estimate a complete (fully-identified) system comprised of (20) substituted into (29) and (49)–(52), subject to (27).\(^\text{14}\)

Assuming that input prices are unobserved and, therefore, generated as latent variables requires stochastic assumptions. Of course, these assumptions have to be consistent with what is known about the sector under study and the nature of sectoral input prices.

Let $p^* = [p^*_l, l = 1, \ldots, L; p^*_j, j = 1, \ldots, J]$. Assume there is a $Q \times 1$ vector of predetermined variables $f_{it}$, so that $E(p^*_t, f_{it}, \zeta_h) = f'_{it}\zeta_h$ for $h = 1, \ldots, J + L$ where

\(^\text{13}\)In a competitive market, the delivered price of coal is a function of the Btu/ton, the percent sulfur/ton, and transportation charges. Given this information, a hedonic regression would yield the implicit price of the percent sulfur/ton. Unfortunately, the DOE publishes data on all these variables except for transportation charges, which are confidential. A way around this would be to obtain data on the mine-mouth price/ton, which could be regressed on Btu/ton and percent sulfur/ton. Again unfortunately, complete mine-mouth data on price/ton linked to a specific mine and utility is confidential.

\(^\text{14}\)One can apply this approach if prices are also missing for elements of $x$ and $y$ by simply generalizing the following procedure.
\(\zeta_h, h = 1, \ldots, J + L\) are \(Q \times 1\) parameter vectors. Define \(F_{il} = I_{J+L} \otimes f_{it}\) so that

\[
p^*_l = F_{il} \zeta + \epsilon_{il}, \quad \epsilon_{il} \sim \mathcal{N}_{J+L}(0, \Sigma_p).
\]

(53)

Thus, the predetermined variables are in \(F_{il}\).

This can be written in the standard multivariate regression form as \(p^* = F\zeta + \epsilon\). For a particular observation we assume

\[
p^* \mid F_{il}, \zeta \sim \mathcal{N}_{J+L}(F_{il} \zeta, \Sigma_{p^*}).
\]

(54)

where \(\bar{\zeta} = 0_{J+L}\) and covariance \(\bar{\Sigma} = 10^4 I_{J+L}\). Relative to the previous approach, we now have to draw from three additional conditional posterior distributions, viz.:

\[
p^* \mid \zeta, \Sigma_{p^*}, \ldots, \zeta \mid p^*, \Sigma_{p^*}, \ldots.
\]

(56)

All three additional conditional posterior distributions are in standard form. Standard results yield the following:

\[
\zeta \mid \cdot \sim \mathcal{N}_{Q'}(\hat{\zeta}, \hat{\Sigma}),
\]

(57)

where \(\hat{\zeta} = \left(F' (I \otimes \Sigma_{p^*}^{-1})^{-1} F\right)^{-1} F' (I \otimes \Sigma_{p^*}^{-1})^{-1} p^*\) and \(\hat{\Sigma} = \left(F' (I \otimes \Sigma_{p^*}^{-1})^{-1} F\right)^{-1}\). For the covariance matrix we have:

\[
p(\Sigma_{p^*} \mid \cdot) \propto |\Sigma_{p^*}|^{-(nT+J+L+1)/2} \exp \left(-\text{tr} \bar{A} \Sigma_{p^*}^{-1}\right),
\]

(58)

where \(\bar{A} = (p^* - F\zeta)'(p^* - F\zeta)\).

Assuming the covariance matrix of errors in (51) and (52) is \(\bar{\Sigma}\) we obtain that

\[
p^*_l \mid \cdot \sim \mathcal{N}_{J+L}(\bar{p}_{il}, \bar{\Sigma}_{p^*}),
\]

(59)

where \(\bar{p}_{il} = (\Sigma_{p^*}^{-1} + \bar{\Sigma}^{-1})^{-1}(\Sigma_{p^*}^{-1} F_{il} \zeta + \bar{\Sigma}^{-1} \bar{G}_{il})\) and \(\bar{\Sigma}_{p^*} = (\Sigma_{p^*}^{-1} + \bar{\Sigma}^{-1})^{-1}\), where \(\bar{G}_{il}\) denotes the RHS of (51) and (52).\(^\text{15}\)

\(^{15}\)Coupled with (27), the price equations in (49)–(52) satisfy the order condition for identification. Of course this discussion is confined to a frequentist view of identification. In Bayesian models, even unidentified parameters can be identified with proper priors. In Table 3 and figures 10 and 11 we provide information on the sensitivity of posterior densities to the choice of priors. 

18
3.3 Measurement of Allocative Inefficiency and its Resource Implications

Using (49)–(52) we can measure allocative inefficiency (AI) as the sum of the error in each input price equation times its corresponding quantity plus the sum of the error in each output price times its corresponding quantity all divided by \( \varrho(p, g) \) (defined in (16)):

\[
AI_{it} = \frac{\sum_{n=1}^{N} \xi_{n,it}x_{n,it} + \sum_{m=1}^{M} \xi_{m,it}y_{m,it} + \sum_{j=1}^{J} \xi_{j,it}\tilde{x}_{j,it} + \sum_{l=1}^{L} \xi_{l,it}\tilde{y}_{l,it}}{\varrho(p, g)}. \tag{60}
\]

To reiterate, this measure depends on the errors from (49)–(52) weighted by corresponding quantities. Since we interpret \( \varrho \) as the change in profits due to an incremental improvement in the production technology, we can decompose \( \varrho \) into two parts: that due to reducing price inefficiency and that due to reducing technical inefficiency. The numerator of \( AI_{it} \) is the former component.

To account for parameter uncertainty, \( AI_{it} \) is averaged across Markov Chain Monte Carlo (MCMC) draws in standard Rao-Blackwell fashion. For each MCMC draw, \( AI_{it} \) in (60) can be transformed as \( \tilde{AI}_{it} := AI_{it} - \min\{AI_{it}\} \). Allocative efficiency (AE) is then computed as

\[
AE_{it} = 100 - \tilde{AI}_{it}. \tag{61}
\]

Since we make this computation for each MCMC draw, the probability of any one firm being fully efficient is very small. Our procedure allows us to resolve the standard problem of relative efficiency estimation by using our variant of the corrected ordinary least-squares technique in a Bayesian framework.

We also compute the estimated percent change in input usage for inputs and outputs due to the firm producing at profit-maximizing levels based on market prices rather than profit-maximizing levels based on shadow prices. For each of equations (49)–(52) we solve the linear system for each draw of parameters and latent variables, which include the estimated value of \( \xi \). A solution \( z^{s,o}_{s,o} \) is obtained for iteration \( s, s = 1, \ldots, S \), for all observations \( o = 1, \ldots, N_T \). This measures the optimal level of inputs subject to market prices, since the solved equations are in terms of these prices. The amounts \( z \) are the current optimal amounts of input usage subject to shadow prices. Then we take the Monte Carlo average \( z^{*}_{s,o} = S^{-1} \sum_{s=1}^{S} z^{s,o}_{s,o} \) to account for parameter uncertainty. Finally, we generate the sample distribution of changes \( c_o \), where \( c_o = \frac{z^{*}_{s,o} - z_o}{z_o} \).
3.4 Posterior Predictive Measure of Cost-Minimization versus Profit Maximization

As indicated above, one can obtain the CM model by estimating the DDF with only the good and bad input price equations. This entails estimation of (20) substituted into (29), (49), and (51), where (27) is imposed during our MCMC iterations.

We wish to be able to compare the CM model to the PM model in terms of their predictive abilities. Suppose all \(T\) observations (say \(y_i \in \mathbb{R}^T\)) of a certain utility \(i \in \{1, \ldots, N\}\) are omitted from the dataset \(Y\) so the new dataset is \(Y_{-i}\). The new posterior is \(p(\theta_i | Y_{-i})\) and the posterior predictive distribution for the \(i\)th utility is:

\[
 p(\tilde{y}_i | Y_{-i}) = \int p(\tilde{y}_i | \theta_i, Y_{-i}) p(\theta_i | Y_{-i}) d\theta_i. \tag{62}
\]

We are interested in the posterior distribution of the absolute prediction error:

\[
 APE = \int_{\mathbb{R}^T} |y_i - \tilde{y}_i|^p(\tilde{y}_i | Y_{-i}) d\tilde{y}_i. \tag{63}
\]

This can be computed with a simple Monte Carlo simulation:

A. Obtain \(\theta_i^s, s = 1, \ldots, S\) using MCMC.

B. Draw \(\tilde{y}_i^s, s = 1, \ldots, S\) from the likelihood, viz., the distribution of \(y_i\) given \(\theta_i^s\).

C. Compute \(APE \simeq S^{-1} \sum_{s=1}^S |y_i - \tilde{y}_i^s|\).

This can be obtained for both the CM and PM models. We use as a predictive measure the ratio:

\[
 R = \frac{APE_{PM}}{APE_{CM}}. \tag{64}
\]

In Fig. 13 we present predictive \(R\) for our sample.

Apparently, the vector \(y_i\) must include only the endogenous variables that are common in the two models. A ratio that has most posterior probability mass \(R < 1\) would indicate that the PM model does a better job in terms of posterior prediction. In step 1, the MCMC is implemented using the same size of burn-in and subsequent draws to convergence as in the original MCMC simulations using the PM and CM model.

3.5 Comparison of PM Models with and without Latent Prices

We compare three shadow-PM models that estimate the DDF together with ancillary equations. All models include the quadratic DDF (20) substituted into (29).
also include in all models two good input price equations containing price inefficiency
terms from (49) for capital and energy (dropping that for labor), and two good output
price equations from (50) for residential and industrial/commercial electricity produc-
tion. With Model I, we ignore endogeneity and simply estimate this system without
instruments. With Model II, we identify the DDF using reduced-form equations for the
bad input and the three bad outputs. In Appendix B we explain in more detail the
reduced-form equations, whose explanatory variables are always the prices of the good
inputs and good outputs, firm dummies, and time dummies.

For Model III we identify the DDF by utilizing the specification for Model II, but
replacing the reduced-form equations with latent price equations from (52) for the two
bad outputs, SO\textsubscript{2} and NO\textsubscript{x}, which are regulated. To determine which variables should
be included in (53), we argue that in equilibrium, the price of each regulated bad output
for the firm should equal its marginal cost of control (which should also equal the price of
an emissions permit if the emission constraint is binding). The marginal cost of control
of each bad output is a function of exogenous prices of inputs used to control that bad
output: the prices of capital, labor, and energy.\textsuperscript{16}

Thus, for Model III, we estimate latent prices for SO\textsubscript{2} and NO\textsubscript{x} as functions of firm
dummies, time dummies, vintage, and the prices of the good inputs, which comprise F\textsubscript{it}
in (53). We do not include a first-order price equation for sulfur, since it is purchased
jointly with the good input, energy, and we lack the data to compute the negative hedonic
price of sulfur as discussed above. Also, we omit the price equation for CO\textsubscript{2}, since it
is an unregulated pollutant with zero price to the firm. We complete this system by
specifying reduced-form equations for the endogenous variables, the quantities of sulfur
and CO\textsubscript{2}. The translation property restrictions for a technology DDF from (27) are
substituted into all equations except for the reduced-form ones. Estimating the DDF in
conjunction with first-order price equations in terms of shadow prices means that the
directions are estimated subject to PM conditions (satisfied with near equality). In this
sense, the estimated directions are optimal. We summarize Bayes factor comparisons of
our models in Fig. 2 discussed below.

We follow the practice of computing and reporting predictive Bayes factors instead
of reporting a single Bayes factor for the entire sample since the possibility always exists
\textsuperscript{16}Our data contains an overall price for each good input, but does not provide separate prices for the
portion used to reduce bad outputs and increase good outputs.
that some observations are particularly influential in driving the Bayes factor in favor of a particular model. For our data, this does not appear to be the case.

4 Econometric Implementation

4.1 Data

The sample consists of an unbalanced panel, subject to attrition, of at most 77 privately-owned electric utilities (whose names are available upon request from the authors) operating in the U.S. over the period 1988-2005, for a total of 1201 observations. This data set is 80% longer than that used by AT, which runs from 1988-1997, thereby omitting a period when many states restructured (deregulated) the generation and delivery of electricity. Approximately 11% of our observations are for periods of restructuring.\textsuperscript{17} A number of firms either merged or sold their assets and dropped out of the sample from 1998 onward. A balanced panel would have yielded 1386 observations.

Since technologies for nuclear, hydroelectric, and internal combustion differ from that of fossil-fuel-based steam generation and because steam generation dominates total production by investor-owned utilities during the time period under investigation, we limit our analysis to this component. We include a full set of 77 firm-specific dummies and omit the intercept in the DDF \textsuperscript{20}.

We model the use of three good inputs (energy, labor, and capital) and one bad input (sulfur) to produce two good outputs (residential and industrial/commercial electricity) and three bad outputs, SO\textsubscript{2}, CO\textsubscript{2}, and NO\textsubscript{x}. From \textit{Federal Energy Regulatory Commission (FERC) Form 1} (1988-2005) we obtain the quantity of energy in total Btu of fuel consumed and the quantity of labor as the number of full-time plus one-half the number of part time employees. In this form, utilities report total capital expenses as the dollars of interest plus depreciation paid by each utility for the sum of production capital and pollution-control capital. From this same form, we decompose total generation into residential and industrial/commercial generation by multiplying total steam output by the percent of sales revenue in each category.

\textsuperscript{17}For a summary of the goals from restructuring the electricity utility industry see Borenstein and Bushnell (2015).
We also calculate prices for the good inputs and good outputs. The price of energy is computed as a weighted average of the cost per million Btu of each fuel, taken from *Department of Energy Information Administration (EIA) Form 767 Boiler Files*. The price of labor is the wage rate, defined as the sum of salaries and wages charged to electric operation and maintenance, divided by the number of full-time plus one-half the number of part-time employees taken from *FERC Form 1* (1988-2005). The price of capital is the yield of the firm’s latest issue of long-term debt adjusted for appreciation and depreciation of capital using the Christensen-Jorgenson (1970) cost of capital formula. These data were taken from *FERC Form 1* (1988-2005) and *Moody’s Public Utility Manual* (1988-2005). The prices of residential and industrial/commercial production are derived as total revenues in each category divided by total sales in that category, where the data are taken from *FERC Form 1* (1988-2005).

Data are available on the quantities, but not prices, for the bad input (sulfur) consumed and all bad outputs generated by the firm. These data are obtained from the *EIA Form 767 Boiler Files*.

We compute \( \tau \) (vintage) using the weighted-average age in years for the firm’s capital, where weights are computed using the firm’s kilowatt-hour (kWh) output, taken from *FERC Form 1* (1988-2005)). This variable and the time dummies are found to be separable from the other inputs and outputs.

In rare cases we encountered missing data for some variables. Whenever necessary we accounted for such data by either using the value of the previous period or the average of the previous and the subsequent period, depending on how related variables changed.

Consumption of total kWh by industrial/commercial customers (66%) was considerably larger than that of residential users (34%) over the sample period. Over the years 1988-1998, total generation remained relatively constant for our sample firms. However, in 1999 total kWh production began a steady decline through the year 2005. For our sample, \( \text{SO}_2/\text{kWh} \) has fallen by about 35% over the sample period in response to the increasingly tight emission restrictions under the \( \text{SO}_2 \) cap-and-trade system. While \( \text{NO}_x/\text{kWh} \) has fallen slightly, \( \text{CO}_2/\text{kWh} \) has risen significantly, since this pollutant is unregulated. All continuous variables are standardized to eliminate variation due to different units of measure when computing directions.
4.2 Satisfying Properties D1-D8

As indicated above, we satisfy the translation property of the DDF, D1, by imposing on each estimated model the restrictions in (27). The restriction D1 implies D2 for our estimated system. To see this, first impose the restrictions in (27) on (25) (with zero on the left-hand-side in order to guarantee frontier efficiency), using \( \vartheta_o = -1, \vartheta = 0 \), and scale \( g \) by \( \lambda \). We obtain

\[
0 = \vec{D}_T(z + \lambda \tilde{\alpha}g; g) = \vec{D}_T(z; g) - \lambda \tilde{\alpha},
\]

which says that the new estimated distance, \( \tilde{\alpha} \), equals \( \lambda^{-1} \) times the original distance, which is a function of the original \( g \). A similar demonstration can be made for the input and output models. We randomly test for concavity, D3, and find that it is satisfied 99% of the time. Non-negativity, D4, is imposed after estimation for all observations via a normalization of the fitted DDF. The monotonicity properties–D5-S, D6-S, D7-W, and D8-W–are satisfied for nearly all of the data via the MCMC estimation process, as explained in the following subsection.

4.3 Statistical Inference in the PM and CM Systems

Our implementation of MCMC relies on a burn-in or transient phase whose length is determined by using Geweke’s diagnostic (1992). In preliminary experiments with various priors, the length of the transient phase ranged from 250,000 to 500,000 iterations. For the baseline prior we used 500,000 burn-in draws followed by another million draws which we use to compute marginal posterior densities and statistics for the functions of interest.

We include a restructuring dummy and also consider the interactions of the restructuring dummy with all first-order terms. Although the restructuring dummy itself is significant with a p-value of .0031, a traditional F-test of the interactions as a whole has a p-value of 0.230. This is really a Bayesian F-test which averages across all MCMC draws.

Monotonicity constraints are often violated in empirical applications. In this paper,
we impose the monotonicity constraints at the means of the variables (which are normalized to zero) and also at a number of points whose distance from the mean is $r$. Since the data are normalized to have unit standard deviation, values of $r$ up to 3 are considered.

Pertaining to all monotonicity properties, the number of violations for the baseline prior is very similar to that for the other priors. Without any restrictions, we have 68 violations. When the restrictions are imposed at the mean we have 31. When the restrictions are imposed at the mean plus a point which is $r = 0.5$ units away from the mean we have 11 violations. Imposing the restrictions with $r = 2$ we have zero violations at the posterior mean of the parameters and a maximum of 2 violations across all MCMC draws. We employ this value for $r$ in the results reported below. The imposition of monotonicity constraints is done using standard rejection techniques.

Our figures and tables pertain only to the shadow-PM system, with the exception of Fig. 13, which indicates that PM is superior to CM. Table 1 summarizes our identification strategies for Models I-III. In Appendix B we explain our use of a Fourier expansion to generate reduced-form expressions used in Models II and III. Based on Bayes factors, increasing the number of terms beyond three provides little improvement. Hence, we employ three terms in our expansion. Figs. 2a and 2b summarize the log of the predictive Bayes Factors against Models I and II and in favor of Model III. Panel 2a (2b) compares the models by omitting utilities (time periods). Clearly Model III is superior based on Bayes factors.

Table 2 reports Bayes factors for different specifications of the productivity equation (30) relative to the linear model, which we employ as model 2 in (38). Throughout the paper, we employ the third-order specification, since it clearly dominates with a Bayes factor of 12.55.  

Figs. 3a and 3b present posteriors of the optimal directions of inputs and outputs for Model III. The mean directions for capital, labor, and residential generation are negative, with posterior means of about -.4, -.65, and -.7, respectively. The mean di-

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Doraszelski, U. and J. Jaumandreu (2013) avoid assuming that all prices are exogenous by omitting subsets of moments involving either lagged wage or lagged price of materials. We have not dropped subsets of moments involving prices since we are assuming that all inputs and outputs are potentially endogenous and we need all prices as instruments. To consider whether lagged prices are better instruments than current period prices, we employ one-period lagged prices for Model III in place of current-year prices. The third-order specification of the productivity equation with current year prices is still superior, in this case with a Bayes factor of 15.66.
rection for energy is positive, with a posterior mean of about .4. Further, the mean
direction for the other good and bad outputs are positive. The largest of these is for
industrial/commercial output with a mean of about .35. The positive mean direction
for energy and the negative mean direction for residential generation are the opposite
signs of what is normally assumed. However, since energy is an input substitute and
industrial/commercial generation is an output substitute, their signs are reasonable. In
addition, the positive directions for industrial/commercial output, NO$_x$, and SO$_2$
emissions are consistent with the assumption that the bads are weakly disposable relative
to industrial/commercial output. Model misspecification would occur if one arbitrarily
assigns directions for bad outputs and all inputs of -1 and directions for good outputs of
+1 or requires that industrial/commercial output and the bad outputs have directions
with opposite signs.

Given estimated optimal-PM directions for measuring distances from the frontier, in
Fig. 4 we present the posterior density of TE for models I-III. Models I and II indicate
mean TE of about .75 and .65, respectively. With Model III, TE is about .85.

Fig. 5 provides for Model III the posterior percentage price distortions (relative
to actual prices) for good inputs and good outputs. The posterior means are small
and negative for all good inputs. From equation (49) the negative price distortions for
inputs indicate that their shadow prices are less than their actual prices. Specifically, the
average shadow price of capital is less than its market price due to super-normal rates
of return allowed in many rate-of-return regulated regimes, which dominate our sample.
This result is consistent with those of other studies of electric utilities referenced above.
From equation (50), somewhat larger positive mean price distortions for good outputs
imply that their shadow prices are slightly greater than their actual prices.

Fig. 6 shows for Model III the posterior percent price distortions (relative to latent
prices) for bad outputs, computed using (52). Mean price distortions are small and
positive for both bad outputs, indicating that shadow prices of the bad outputs exceed
their market prices.

In Fig. 7, for Models I-III we provide the posterior densities of AE computed from
equation (61) for all inputs and outputs. Mean AE for Models I-III is about .85, .66,
and .64, respectively.

In Fig. 8 we provide for Model III the posteriors of the percent changes in the usage
of inputs and outputs if firms were required to produce at profit-maximizing levels based
on actual (market) prices rather than profit-maximizing levels based on shadow prices. We use the methodology described in section 3.3 above. The computed percent change for a given input or output is a function of marginal rates of transformation and price changes of all other inputs and outputs, through the solution of a system of equations. Thus, the percent changes in usage are not linearly related to individual price distortions. The mean posterior changes in usage are -.04 and -.07 for capital and labor, respectively, and .05 for energy. The mean posterior changes for good and bad outputs range from .02 to .04. All of these distortions are relatively small.

Fig. 9 provides posterior densities for RTS, PC, TC, and EC for Model III. The posterior mean of RTS is about .94, indicating slightly decreasing returns to scale. The posterior mean of EC is slightly negative and its posterior distribution is somewhat less disperse than are those of PC and TC. The posterior mean of TC is slightly greater than .01 and the posterior mean of PC is slightly less than .01.

Table 3 indicates the range in 10,000 priors used for posterior sensitivity analysis. Fig. 10 reports the resulting posteriors for inputs and outputs for Model III. We focus on changes in the posteriors of inputs and outputs relative to the baseline prior. To minimize computational costs of this posterior comparison for each of the 10,000 different priors, we use Sampling-Importance-Resampling (SIR) following Smith and Gelfand (1992) and Rubin (1987). Given a set of posterior draws \( \{\beta^{(s)}, s = 1, ..., S\} \) for a model with a given baseline prior, say \( p_0(\beta) \), approximate draws from the same model with an alternative prior \( p(\beta) \) can be obtained using the SIR algorithm. This attaches weights, \( Y_s = \frac{p(\beta^{(s)})}{p_0(\beta^{(s)})} \), to the original draws and resampling is used with these normalized weights avoiding the reuse of MCMC\(^{19}\). All sensitivities are quite small.

Fig. 11 reports the sensitivity of changes in posterior means for structural parameters of Models I-III to 10,000 different priors. Since the use of SIR does not require new MCMC computations, it is particularly suited to large-scale prior sensitivity analysis as in our case. The 10,000 different priors are generated from the baseline prior of each parameter or block using the hyperparameters of these priors. If the vector of hyperparameters is collectively denoted by \( \delta \in \mathbb{R}^\Delta \) new priors are generated using \( \delta^* = \delta + \kappa \) where \( \kappa \) is uniformly distributed in \([-B_1, B_2]^\Delta\).

We set \( B_1 = B_2 = 10 \) for hyperparameters that can be defined over the real line and \( B_1 = 0.001, B_2 = 10 \) for positive hyperparameters. Performing the same sensitivity

\(^{19}\)The size of the resample is set to 20% of the number of available MCMC draws.
analysis experiment when only the priors for the structural parameters are allowed to change, the changes in posterior means are modest, suggesting that posterior MCMC is quite robust.

After estimating the shadow PM model, we compute for Model III the posterior means of relative shadow prices over time for bad outputs from equation (52). As reported in Fig. 12, these prices all decline over time, consistent with historically declining costs of pollution control. The relative prices of \( \text{SO}_2 \) and \( \text{NO}_x \) are consistent with estimates of control costs from the Integrated Environmental Control Model (Rubin, 2009).

In order to compare the predictive accuracy of the shadow-CM and shadow-PM models, we compute the marginal posterior of the predictive measures of \( R \). For five randomly chosen utilities, provided in Fig. 13, \( R \) values range from zero to .5. The mean values range from approximately .01 to about .1, indicating that the shadow-PM model is strongly preferred to the shadow-CM model.

Finally, in Figs. 14-15 and Table 4 we report for Model III the partial elasticities of \( \omega \) in (30) with respect to \( \omega_{t-1}, p_x, p_y, \pi, \tau \) and \( t \). When implementing (32)-(35) we do not impose monotonicity of profits with respect to \( \omega \). However, we find that the required monotonicity is satisfied for 99% of our observations.

Lagged \( \omega \) is the most important variable affecting \( \omega \) with a positive elasticity of .43. Reducing \( p_E \) is the second most important variable with an elasticity of -.41. Reductions in \( p_K \) and \( p_L \) are more important than an increase in the prices of residential and industrial production. Profits have a very small but positive effect on productivity. All elasticities except for those of \( t \) and \( \tau \) are significant at the .05 level using a two-tailed test, indicating little affect of time itself or the aging of capital stock.

5 Conclusions

Using a Bayesian approach, our contributions to the productivity literature are fourfold. First, we estimate unique optimal-PM directions for each good input, each good output, and all regulated bad outputs. Second, we allow the potential endogeneity of all inputs and outputs. This entails identifying the DDF by assuming a data generating process for the latent prices as instruments to replace missing prices for the regulated bad outputs. Then we estimate the DDF jointly with the first-order price equations derived from PM for good inputs, good outputs, and two controlled pollutants. The bad input,
sulfur, and the uncontrolled pollutant, CO₂, are identified using reduced-form equations. We also estimate the corresponding CM model. Third, we avoid the typical assumption that firms respond to actual prices by estimating firm-specific shadow prices. Fourth, we generalize and provide an alternative to the approaches of OP (1996) and LP (2003). We accomplish this by treating all input and output quantities as potentially endogenous and deriving productivity as a function of lagged productivity, profits, vintage, time, and the prices of good inputs and outputs. From this function, we compute TC, EC, PC, and partial elasticities.

Using MCMC methods, we generate posterior densities for the parameters and latent variables of our system using an unbalanced panel of 77 U.S. electric utilities for the years 1988-2005. Using Bayesian criteria, the shadow PM model is superior to the shadow CM model in terms of predictive ability. Optimal directions subject to shadow PM differ from their typically assumed values (-1 for all bad outputs and all inputs, and +1 for good outputs). Estimated price distortions for the good inputs indicate that efficient levels of usage are slightly lower for capital and labor but are moderately higher for energy. Mean PC is slightly less than 0.01. The major factors augmenting productivity are an increase in lagged productivity and a reduction in energy prices. Changes in posterior means of the structural parameters and the directions are highly insensitive to a wide range of different priors.

References


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Caves, D.W., Christensen, L. R., Diewert, W. E., 1982b, Multilateral comparisons of output, input, and productivity using superlative indexes. The Economic Journal 92, 73-86.


Rubin, E. S., 2009, Integrated Environmental Control Module (IECM), version 6.2.4, (developed under contracts from DOE), Carnegie Mellon University.


### 6 Tables and Figures

Table 1: Identification Strategies for Models I-III

<table>
<thead>
<tr>
<th>Model</th>
<th>Strategy</th>
<th>Variables with Identifying Equations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Model I</td>
<td>Ignore Endogeneity</td>
<td>none</td>
</tr>
<tr>
<td>Model II</td>
<td>Add Reduced-Form Equations</td>
<td>$\tilde{x}_S$ and $\tilde{y}$</td>
</tr>
<tr>
<td>Model III</td>
<td>Add Reduced-Form Equations</td>
<td>$\tilde{x}<em>S$, $\tilde{y}</em>{CO_2}$</td>
</tr>
<tr>
<td></td>
<td>Add Latent-Price Equations</td>
<td>$\tilde{p}<em>{SO_2}$, $\tilde{p}</em>{NO_x}$</td>
</tr>
</tbody>
</table>
Table 2. Bayes factors for different specifications of productivity equation

<table>
<thead>
<tr>
<th>Specification</th>
<th>BF</th>
</tr>
</thead>
<tbody>
<tr>
<td>linear</td>
<td>1.000</td>
</tr>
<tr>
<td>second-order</td>
<td>5.301</td>
</tr>
<tr>
<td>third-order</td>
<td>12.545</td>
</tr>
<tr>
<td>fourth-order</td>
<td>2.271</td>
</tr>
<tr>
<td>NN, $G = 1$</td>
<td>6.412</td>
</tr>
<tr>
<td>NN, $G = 2$</td>
<td>5.812</td>
</tr>
<tr>
<td>NN, $G = 3$</td>
<td>4.023</td>
</tr>
<tr>
<td>NN, $G = 4$</td>
<td>1.167</td>
</tr>
<tr>
<td>Fourier, unrestricted $\mathbf{A}$</td>
<td>2.519</td>
</tr>
</tbody>
</table>

Notes: The linear model is obtained from (33) by setting $\mathbf{A} = 0$, “NN” stands for “neural network” and the specification is given by (32). The Fourier approximation is as in (34). The model “Fourier, unrestricted $\mathbf{A}$” corresponds to (34) with an unrestricted $\mathbf{A}$ matrix.
### Table 3. Baseline priors and range of variation for parameters to generate the 10,000 priors for posterior sensitivity analysis

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Equation</th>
<th>Value</th>
<th>Range of variation</th>
</tr>
</thead>
<tbody>
<tr>
<td>$g$</td>
<td>Appendix (C.1.5)</td>
<td>Semi-informative form</td>
<td>fixed</td>
</tr>
<tr>
<td>$c$</td>
<td>(27)</td>
<td>$10^{-4}$</td>
<td>$10^{-3}$ to $10^{-8}$</td>
</tr>
<tr>
<td>$\bar{g}$</td>
<td>(39)</td>
<td>-1 or +1</td>
<td>fixed</td>
</tr>
<tr>
<td>$\Sigma_g$</td>
<td>(39)</td>
<td>Wishart($\nu, A$)</td>
<td>See note below</td>
</tr>
<tr>
<td>$\Delta$</td>
<td>Appendix (B.1)</td>
<td>Wishart($\nu, A$)</td>
<td>See note below</td>
</tr>
<tr>
<td>$\pi$</td>
<td>Appendix (B.1)</td>
<td>$10^4$ varies from 10 to $10^9$</td>
<td>$10^{-2}$ varies from $10^{-4}$ to 1</td>
</tr>
<tr>
<td>$\sigma_u$</td>
<td>Appendix (C.2.5.1)</td>
<td>$\frac{Q}{\sigma_u^2} \sim \chi^2_{\nu}, \ Q = 0.1, \ \nu = 1$</td>
<td>$Q \in [10^{-3}, 100], \ \nu \in [0.01, 20]$</td>
</tr>
<tr>
<td>$F$</td>
<td>Appendix (B.2)</td>
<td>5</td>
<td>Chosen initially using BIC</td>
</tr>
</tbody>
</table>

Note: Wishart priors are of the form $p(\Sigma) \propto |\Sigma|^{(\nu-m-1)/2} \exp \left( tr(A^{-1}\Sigma) \right)$ where the dimensionality of the matrix is $m \times m$ generically, and $\nu, A$ are prior parameters. We set the parameter $\nu$ equal to 0.1 times the sample size and $A^{-1} = 0.001I$. 


Table 4. Partial Elasticity of $\omega_{it}$ with respect to $p_x, p_y, \pi, \tau, t$

<table>
<thead>
<tr>
<th>Variable</th>
<th>Partial Elasticity</th>
<th>Standard Error</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\omega_{i,t-1}$</td>
<td>0.432</td>
<td>(0.081)</td>
</tr>
<tr>
<td>$p_E$</td>
<td>-0.414</td>
<td>(0.122)</td>
</tr>
<tr>
<td>$p_L$</td>
<td>-0.235</td>
<td>(0.092)</td>
</tr>
<tr>
<td>$p_K$</td>
<td>-0.167</td>
<td>(0.072)</td>
</tr>
<tr>
<td>$p_{RES}$</td>
<td>0.045</td>
<td>(0.023)</td>
</tr>
<tr>
<td>$p_{IND}$</td>
<td>0.071</td>
<td>(0.022)</td>
</tr>
<tr>
<td>$\pi_{it}$</td>
<td>0.044</td>
<td>(0.011)</td>
</tr>
<tr>
<td>$t$</td>
<td>0.0014</td>
<td>(0.0013)</td>
</tr>
<tr>
<td>$\tau_{it}$</td>
<td>-0.157</td>
<td>(0.032)</td>
</tr>
</tbody>
</table>
Fig. 1: Movement from Interior Point to Profit-Maximizing Position
Fig. 2: Log of Predictive Bayes Factors

(a) Log predictive Bayes factors in favor of endog. directions (omitting utilities)

(b) Log predictive Bayes factors in favor of endog. directions (omitting time periods)
Fig. 3: Posterior Densities for Directions
Fig. 4: Posterior Density for Technical Efficiency for Models I-III
Fig. 5: Posterior Densities for Price Distortions—Good Inputs and Good Outputs
Fig. 6: Posterior Densities for Price Distortions–Bad Outputs

c. Post. Distributions of price distortions

\[ \text{density} \]

\[ \text{distortion} \]
Fig. 7: Posterior Density for Allocative Efficiency for Models I-III
Fig. 8: Posterior Densities of Change in Inputs and Outputs
Fig. 9: Posterior Densities for RTS, PC, TC, and EC
Fig. 10: Posterior Densities for Sensitivity of Changes in Inputs and Outputs to 10,000 Priors
Fig. 11: Posterior Densities for Sensitivity of Changes in Structural Parameters to 10,000 Priors
Fig. 12: Posterior Means of Relative Prices of Bad Outputs over Time from PM Model
Fig. 13: Posterior Density of Predictive Measure, R, for PM Model
Fig. 14: Partial Elasticities for $\omega_{ij}$ for the PM Model
Fig. 15: Partial Elasticities for $\omega_{it}$ for the PM Model
Appendices

A.1: Monotonicity for Good Inputs

In what follows we suppress the arguments \((ω, τ, t)\) in equations (1) and (2) for \(T\) and \(\vec{D}_T\) for the sake of simplicity.

For \((x, \tilde{x}, y, \tilde{y}) \in T, x' \geq x \Rightarrow (x', \tilde{x}, y, \tilde{y}) \in T\). Let \(\beta^*\) be the solution to (2). Therefore, it is optimal and feasible in (2). Then

\[(x + \beta^* g_x, \tilde{x} + \beta^* g_{\tilde{x}}, y + \beta^* g_y, \tilde{y} + \beta^* g_{\tilde{y}}) \in T.\]

Further, let

\[\vec{D}_T(x', \tilde{x}, y, \tilde{y}; g) = \beta' = \sup\{\beta : (x' + \beta g_x, \tilde{x} + \beta g_{\tilde{x}}, y + \beta g_y, \tilde{y} + \beta g_{\tilde{y}}) \in T\}.\]

(A.1.1)

Assume that \(x' \geq x\). Then

\[(x' + \beta^* g_x, \tilde{x} + \beta^* g_{\tilde{x}}, y + \beta^* g_y, \tilde{y} + \beta^* g_{\tilde{y}}) \geq (x + \beta^* g_x, \tilde{x} + \beta^* g_{\tilde{x}}, y + \beta^* g_y, \tilde{y} + \beta^* g_{\tilde{y}}).\]

Since inputs are strongly disposable,

\[(x' + \beta^* g_x, \tilde{x} + \beta^* g_{\tilde{x}}, y + \beta^* g_y, \tilde{y} + \beta^* g_{\tilde{y}}) \in T.\]

Hence \(\beta^*\) is feasible in (A.1.1) although not necessarily optimal in (A.1.1). Thus,

\[\beta' = \vec{D}_T(x', \tilde{x}, y, \tilde{y}; g) \geq \beta^* = \vec{D}_T(x, \tilde{x}, y, \tilde{y}; g).\]

We conclude that

\[x' \geq x \Rightarrow \vec{D}_T(x', \tilde{x}, y, \tilde{y}; g) \geq \vec{D}_T(x, \tilde{x}, y, \tilde{y}; g).\]

Now to convert this into a partial derivative, let \(x' = x + h\), where \(h > 0\). Then \(x' \geq x\) implies that

\[\vec{D}_T(x + h, \tilde{x}, y, \tilde{y}; g) \geq \vec{D}_T(x, \tilde{x}, y, \tilde{y}; g)\]

or

\[\frac{\vec{D}_T(x + h, \tilde{x}, y, \tilde{y}; g) - \vec{D}_T(x, \tilde{x}, y, \tilde{y}; g)}{h} \geq 0.\]

Then, since

\[\frac{\partial \vec{D}_T(x, \tilde{x}, y, \tilde{y}; g)}{\partial x_n} = \lim_{h \to 0} \frac{\vec{D}_T(x + h, \tilde{x}, y, \tilde{y}; g) - \vec{D}_T(x, \tilde{x}, y, \tilde{y}; g)}{h},\]

\(n = 1, \ldots, N,\)
we obtain
\[ \partial \hat{D}_T(x, \tilde{x}, y, \tilde{y}; g)/\partial x_n \geq 0, n = 1, \ldots, N. \] (A.1.2)

**A.2: Monotonicity for Good Outputs**

Assuming the strong disposability of good outputs in D6–S,
\[ (x, \tilde{x}, y, \tilde{y}) \in T, y' \leq y \Rightarrow (x, \tilde{x}, y', \tilde{y}) \in T. \] (A.2.1)

In a similar manner with D5–S, assume that \( y' \leq y \) and let \( \beta^* = \hat{D}_T(x, \tilde{x}, y, \tilde{y}; g) \) be the solution to (2). Then
\[ (x + \beta^* g_x, \tilde{x} + \beta^* g_{\tilde{x}}, y' + \beta^* g_y, \tilde{y} + \beta^* g_{\tilde{y}}) \in T. \]

Since outputs are strongly disposable, we have
\[ (x + \beta^* g_x, \tilde{x} + \beta^* g_{\tilde{x}}, y' + \beta^* g_y, \tilde{y} + \beta^* g_{\tilde{y}} \in T). \]

Thus,
\[ \hat{D}_T(x, \tilde{x}, y', \tilde{y}; g) = \beta^* \geq \hat{D}_T(x, \tilde{x}, y, \tilde{y}; g) = \beta^*. \]

We conclude that
\[ y' \leq y \Rightarrow \hat{D}_T(x, \tilde{x}, y', \tilde{y}; g) \geq \hat{D}_T(x, \tilde{x}, y, \tilde{y}; g) \]

or that (following the steps in D5 to obtain a partial derivative)
\[ \partial \hat{D}_T(x, \tilde{x}, y, \tilde{y}; g)/\partial y_m \leq 0, m = 1, \ldots, M. \] (A.2.2)

Note that D5–S and D6–S do not depend on the signs of the elements of \( g \).

**B: Reduced-Form Equations for Endogenous Bads**

Suppose \( \tilde{z} = [\tilde{x}', \tilde{y}']' \), which is a \( (J + L) \times 1 \) vector, denotes these endogenous variables, and \( W \) denotes the \( (J + L) \times K \) matrix of \( K \) weakly exogenous or predetermined variables in the model, including relative prices or their logs. Let \( \pi \) denote a \( K \times 1 \) vector of coefficients, and \( \eta \) a \( (J + L) \times 1 \) vector of error terms such that \( \eta \sim \mathcal{N}_{J+L}(0, \Delta) \):
\[ \tilde{z} = W \pi + \eta, \] (B.1)

where \( \Delta \) is to be estimated. Equation (B.1) can be used to complete the system of (29), (49), and (50), providing equations in reduced form for the endogenous variables \( \tilde{x} \) and \( \tilde{y} \), for which price data are not available. Matrix \( W \) can be used so that (B.1) is, in fact, a semi-parametric formulation, contrary to standard LIML, where a linear reduced form
is used. This treatment has been absent in most Bayesian treatments of LIML. Given a basic set of weakly exogenous or predetermined variables, a Fourier expansion may be used (which amounts to including in \( W \) trigonometric terms of properly transformed variables in the data set) and the order of the expansion can be used to control the flexibility of the approximation. We reiterate that for any \( \chi \in \mathbb{R} \) an unknown univariate function, \( \varsigma : [-\pi, \pi] \to \mathbb{R} \) can be approximated as follows:

\[
\varsigma (\chi) \cong \alpha_0 + \sum_{f=1}^{F} \alpha_f \cos (fz) + \sum_{f=F+1}^{2F} \alpha_f \sin ((f - F)z),
\]

where \( z = 2\pi (\chi - \chi_{\min}) / (\chi_{\max} - \chi_{\min}) - \pi \), and \( F \in \{1, 2, \ldots\} \) denotes the order of the approximation. For a multivariate expansion we take the sum of univariate expansions (so as to economize somewhat on the number of parameters). For \( K \) variables in the basic set, Fourier expansions of common order \( F \) imply that we have \( 2FK \) parameters like \( \alpha_f \) (omitting, of course, the constant terms which are included separately in (B.2)).

If the variables in \( W \) are denoted by \( \varpi_1, \varpi_2, \ldots, \varpi_{J+L} \) the idea is to use (B.2) and define new variables:

\[
\tilde{\varpi}_{nf}^{(1)} = \cos (fz_n), \quad \tilde{\varpi}_{nf}^{(2)} = \sin (f\varpi_n), \quad n = 1, \ldots, J+L, \quad f = 1, \ldots, F,
\]

where \( z_n = 2\pi (\varpi_n - \varpi_{n,\min}) / (\varpi_{n,\max} - \varpi_{n,\min}) - \pi, \quad n = 1, \ldots, J+L. \) Then we define the new matrix:

\[
W^* = [W, \tilde{W}^{(1)}, \tilde{W}^{(2)}],
\]

in obvious notation whose dimension is \((J+L) \times K'\) and \( K' = 1 + (1 + 2F)K \) where the first 1 stands for the intercept, followed by the \( K \) variables in the basic data set \( W \), and then followed by the \( 2FK \) trigonometric elements of the Fourier approximation. The reduced form in (B.1) is then replaced by:

\[
\tilde{z} = W^* \pi + \varepsilon.
\]

To control for the proliferation of parameters\(^\text{20}\) we use a special prior:

\[
\pi_{1:K+1} \sim \mathcal{N}_{K+1} (0, 10^3 I_K),
\]

\[
\pi_{K+2:2KF} \sim \mathcal{N}_{K(2F-1)} (0, 10^{-2} \cdot I_{K(2F-1)}).
\]

The first \( K + 1 \) parameters (corresponding to the intercept and the \( K \) variables in the basic set \( W \)) have diffuse normal priors. The coefficients corresponding to the Fourier terms are relatively tightly concentrated around zero to reflect the prior opinion that a linear expansion is likely to be best. Of course, in light of the data, such prior beliefs

---

\(^{20}\)With \( K = 10 \) basic variables and an order \( F = 5 \) we would end up with 100 trigonometric terms.
can be updated.

C: Bayesian Analysis

C.1: Priors and Posteriors

We use (29) to express the quadratic technology DDF as

\[ 0 = \gamma' \delta_{it} + \frac{1}{2} \delta_{it}' \Gamma \delta_{it} + v_{it} + \omega_{it} - u_{it}, \tag{C.1.1} \]

where \( \gamma = [\gamma_w, w = 1, ..., W]' \), \( \Gamma = [\gamma_{ww}'] \), \( i = 1, ..., N, t = 1, ..., T \), and we set \( \omega_{it} = 0 \). To economize on notation we define \( \gamma = [\gamma', \text{vec}(\Gamma)']' \), the structural parameters of the system.

Combining (C.1.1) with equations (49), (50), and the reduced form equations in (B.5), we have a simultaneous equations model whose Jacobian of transformation is the following:

\[ J(g_i, \xi_{it}, \gamma) = \begin{bmatrix} -\gamma - \Gamma \delta_{it} \\ \pi_{it}' (g_i, \xi_{it}) \end{bmatrix} \tag{C.1.2} \]

where \( \pi_{it}' = [1_N, -1_M]' \) and

\[ \pi_{it}' (g_i, \xi_{it}) = \sum_{m=1}^{M} p_{m, it} g_{m, i} - \sum_{n=1}^{N} p_{n, it} g_{n, i} + \sum_{m=1}^{M} \xi_{m, it} g_{m, i} - \sum_{n=1}^{N} \xi_{n, it} g_{n, i}. \]

Since we have introduced optimization error terms \( v_n \) and \( v_m \) in (49) and (50), the price distortions \( \xi_{it} \) will be viewed as nonlinear random effects in the system. For stronger identification we assume that the vector of directions \( g_i \) is time-invariant, which is a reasonable assumption in our context.

In the system consisting of (C.1.1), (49), and (50), the prior for the standard noise component is as follows:

\[ V_{it} = [v_{it}, v_{N, it}', v_{M, it}']' \sim \mathcal{N}_{N'+M+1} (0, \Sigma), \tag{C.1.3} \]

where \( v_{N'} = [v_{n, it}, n = 1, ..., N']' \), \( v_M = [v_{m, it}, m = 1, ..., M]' \), and \( N' = N - 1 \) (since one input price equation is omitted). Here, \( v_{it} = -\sum_{w=1}^{W} \gamma_w z_w - \sum_{w=1}^{W} \sum_{w'=1}^{W} \gamma_{ww'} z_w z_{w'} + u_{it} \) and similarly for \( v_{n, it} \) and \( v_{m, it}.\)

Due to the constraints in (27), it is difficult to place informative priors on both \( \gamma \) and \( g_i \). One problem is that the directions are firm-specific but the structural parameters are not, so these restrictions cannot hold exactly for each observation. The problem would, of course, disappear under the assumption that all firms have the same direction vectors.

\[ ^{21} \text{This implies that the } v's \text{ are now considered as functions of the data and parameters to economize on notation.} \]
However, we feel that this assumption is too restrictive. To overcome the problem we write our set of restrictions in the form

\[ F(\gamma, g_i) = A(\gamma) g_i = \vartheta_i, \quad i = 1, \ldots, n, \quad (C.1.4) \]

where \( F \) is a vector function in \( \mathbb{R}^{M+L+N+J+1} \). Here, \( \vartheta_i \sim \mathcal{N}_{M+L+N+J+1}(a, \sigma^2 I) \) where the first element of \( a \) is \( -1 \) and the remaining elements are zero (see discussion below (27)). The function \( F \) is nonlinear as it involves products of \( \gamma \) and \( g_i \).

The conditional prior distribution for \( \gamma \) given \( g \) is then:

\[ p(\gamma | g) = \prod_{i=1}^{N} \exp \left( -\frac{1}{2} g_i' A(\gamma)' A(\gamma) g_i \right) . \quad (C.1.5) \]

Given our notation the joint posterior distribution (augmented by all latent variables) can be written as follows:

\[ p(\gamma, \pi, \xi, g, u, \Sigma, \Omega, \Delta | \mathcal{Y}) \propto \left| \Sigma \right|^{NT(N+M)} \exp \left( -\frac{1}{2} \sum_{i=1}^{N} \sum_{t=1}^{T} V_{it}' \Sigma^{-1} V_{it} \right) \]

\[ \left| \Omega \right|^{-\frac{T(N'+M')}{2}} \exp \left( -\frac{1}{2} \sum_{i=1}^{N'} \sum_{t=1}^{T} \xi_{it}' \Omega^{-1} \xi_{it} \right) \]

\[ |\Delta|^{-\frac{NT(J+L)}{2}} \exp \left[ -\frac{1}{2} \sum_{i=1}^{N} \sum_{t=1}^{T} (\tilde{z}_{it} - W_{it} \pi)' \Delta^{-1} (\tilde{z}_{it} - W_{it} \pi) \right] \prod_{i=1}^{N} \prod_{t=1}^{T} J(g_i, \xi_{it}, \gamma) \cdot p(u, \xi, \Sigma, \Omega, \Delta, \pi) \cdot p(\gamma | g) p(g), \quad (C.1.6) \]

where we set \( \omega_{it} = 0 \). Lines 2-4 of the joint posterior are the contributions to the likelihood. On the last line, the second from the last term denotes the prior for the one-sided error term \( u \), the price distortions \( \xi \), various covariance matrices of the model, and the reduced-form parameters, \( \pi \), all of which are assumed to be independent. The last two terms denote the prior on the structural parameters \( \gamma \) and the directions \( g_i \), which we denote collectively by \( g \). The data is collectively denoted by \( \mathcal{Y} \). See Table 2 for the values of all priors and ranges of variation actually employed.

Combined with the rest of the posterior in (C.1.6), (C.1.5) acts as a ‘penalty function’ whenever there is a deviation of \( F \) from its prescribed values in (27). The value of \( c \) is chosen a priori.

For all covariance matrices (denoted generically by \( \Phi \)) the prior has the inverted Wishart form:

\[ p(\Phi | \nu_{\Phi}, A_{\Phi}) \propto |\Phi|^{-\nu_{\Phi}+m_{\Phi}+1} \exp \left( -\frac{1}{2} tr[A_{\Phi} \Phi^{-1}] \right) \quad (C.1.7) \]

where \( \nu_{\Phi} \) is the degrees of freedom, \( m_{\Phi} \) is the dimensionality of \( \Phi \), and where \( A_{\Phi} \) is a positive definite matrix of coefficients that determine other features of the distribution,
including its location.

We can easily add bad inputs and bad outputs by including their equations, (51) and (52). In this case, we must return to the beginning of this subsection and generalize \( J \), \( \pi_{it}^* \), \( V_{it} \), and the joint posterior distribution to accommodate \( J \) bad inputs and \( L \) bad outputs.

C.2: Markov Chain Monte Carlo Schemes

C.2.1: Drawing structural parameters \( \gamma \)

From (C.1.6) or the formulation in (20) and (29) along with (49), (50), and (B.5), a complication arises from the presence of the \( \gamma \) parameters in the Jacobian in (C.1.2) given all the other parameters. The problem is the same with parameters like \( g_i \) and \( \xi_{it} \) which also appear in the Jacobian.

Assuming that we have \( G \) equations in our system, equations (20) and (29) along with (49) and (50) can be written in the general form:

\[
Y = X\alpha(\gamma) + V - u
\]  

where \( Y \) and \( V \) are \((NTG \times 1)\) vectors denoting the data for left-hand-side variable in (20) and (29) along with (49) and (50), conditional on all other parameters (particularly parameters like \( g_i \) and \( \xi_{it} \), and the \((NTG \times 1)\) vector \( u = (u_1, 0_{NT(G-1)})' \), where \( u_1 = [u_{it}] \) is a \((1 \times NT)\) vector of \( u_{it}, i = \ldots, N; t = 1, \ldots, T \) and \( 0_{NT(G-1)} \) is a \((1 \times NT(G-1))\) vector of zeroes. The notation \( \alpha(\gamma) \) means that there are cross-equation restrictions among the parameters \( \gamma \) arising from the symmetry conditions and from (27). These appear in the DDF and in the first-order conditions. For example, (49) and (50) are conditioned on \( g_i \) and \( \xi_{it} \) in a standard SURE system. The matrix of explanatory variables \( X \), has \( NTG \) columns and number of rows equal to the number of explanatory variables in all the equations. We write the system as

\[
\begin{bmatrix}
Y_1 \\
Y_2 \\
\vdots \\
Y_G \\
\end{bmatrix} =
\begin{bmatrix}
X_1 \\
X_2 \\
\vdots \\
X_G \\
\end{bmatrix}
\alpha(\gamma) +
\begin{bmatrix}
V_1 - u_1 \\
V_2 \\
\vdots \\
V_G \\
\end{bmatrix}
\]  

where \( Y_g \) \((g = 1, \ldots, G)\) denotes the left-hand-side variable in the \( g^{th} \) equation \((G = N + M)\), \( X_1 \) is the matrix of regressors in the distance function, and \( X_g \) is the matrix of regressors in the remaining price equations (49) and (50). The coefficient vector \( \alpha(\gamma) \) is essentially \( \alpha(\gamma) = S\gamma \) where \( S \) is a selection matrix. The model can be written as
\[ Y = X\gamma + V - \begin{bmatrix} u_1 \\ 0_{N_T(G-1)} \end{bmatrix} \quad (C.2.1.3) \]

if we redefine \( X \) properly. For example \( X_1 \) contains regressors of the technology DDF, and \( X_2, ..., X_G \) contain the corresponding elements given by the price equations in (49) and (50), where the symmetry restrictions and (27) have been imposed.

Since \( \mathcal{E}(V'V) = \Sigma \otimes I_{N_T} \) it is easy to see that the conditional distribution of \( \alpha \) is

\[
\alpha|\Sigma, ..., \mathcal{Y} \sim N \left( \left[ X' \left( \Sigma^{-1} \otimes I \right)^{-1} X \right]^{-1} \left[ X' \left( \Sigma^{-1} \otimes I \right)^{-1} Y \right], \left[ X' \left( \Sigma^{-1} \otimes I \right)^{-1} X \right]^{-1} \right) \quad (C.2.1.4)
\]

apart from the Jacobian term. In this conditional posterior we also have to account for the presence of \( \alpha \) in the conditional prior in (C.1.5). The posterior conditional distribution of \( \Sigma^{-1} \) is, however, in the Wishart family, so that

\[
p(\Sigma^{-1}|\alpha, ..., \mathcal{Y}) \propto |\Sigma^{-1}|^{(\nu + N_T - (G+1))/2} \exp \left( -\frac{1}{2} tr \left[ A_\Sigma \Sigma^{-1} \right] \right). \quad (C.2.1.5)
\]

Turning attention to \( \gamma \), it also appears in the Jacobian so its complete posterior is the product of the normal density corresponding to (C.2.1.3) and the Jacobian terms in (C.1.6).

There are two solutions to this problem:

(a) We use (C.2.1.4) as a candidate generating function and then we use a Metropolis-Hastings update to maintain the correct posterior conditional distribution.

(b) We compute the gradient and Hessian of the full conditional kernel posterior distribution. Then we apply a Girolami-Calderhead (2011) (GC) update for \( \gamma \), see below.

A Metropolis-Hastings update for a block of parameters, say \( \vartheta \), when the proposal is \( \mathcal{N}(\hat{\vartheta}, \hat{V}) \) whose density we denote by \( f_N(\vartheta; \hat{\vartheta}, \hat{V}) \) is done as follows. Given that the current MCMC draw is \( \vartheta^o \) and the candidate vector from the proposal is \( \vartheta^c \), the candidate is accepted with the Metropolis-Hastings probability

\[
\min \left\{ 1, \frac{p(\vartheta^c|..., \mathcal{Y})}{p(\vartheta^o|..., \mathcal{Y})} \cdot \frac{f_N(\vartheta^o; \hat{\vartheta}, \hat{V})}{f_N(\vartheta^c; \hat{\vartheta}, \hat{V})} \right\}
\]

The GC procedure is as follows. Suppose \( \mathcal{L}(\beta) = \log p(\beta|\Sigma, u, \Lambda, \mathcal{Y}) \) is used to denote for simplicity the log posterior of any block of parameters \( \beta \). Moreover, define

\[
G(\beta) = \text{est. cov.} \frac{\partial}{\partial \beta} \log p(\mathcal{Y}|\beta, \Sigma, u, \Lambda)
\]
the empirical counterpart of
\[ G_o (\beta) = -E_{\mathcal{Y} | \beta} \frac{\partial^2}{\partial \beta \partial \mathcal{Y}} \log p (\mathcal{Y} | \beta, \Sigma, u, \Lambda). \]

The Langevin diffusion is given by the following stochastic differential equation:
\[ d\beta (t) = \frac{1}{2} \tilde{\nabla}_{\beta} \mathcal{L} \{ \beta (t) \} \, dt + dB (t) \]
where
\[ \tilde{\nabla}_{\beta} \mathcal{L} \{ \beta (t) \} = -G^{-1} \{ \beta (t) \} \cdot \tilde{\nabla}_{\beta} \mathcal{L} \{ \beta (t) \} \]
is the so-called “natural gradient” of the Riemann manifold generated by the log posterior. The elements of the Brownian motion are
\[ G^{-1} \{ \beta (t) \} dB_i (t) = | G \{ \beta (t) \} |^{-1/2} \sum_{j=1}^{K_{\beta}} \left[ G^{-1} \{ \beta (t) \} \frac{\partial G (\beta (t))}{\partial j} \right] dt + \left[ \sqrt{G \{ \beta (t) \}} dB (t) \right]. \]

The discrete form of the stochastic differential equation provides a proposal as follows:
\[ \tilde{\beta}_i = \beta_i^o + \frac{\varepsilon^2}{2} \left\{ G^{-1} (\beta^o) \nabla_{\beta} \mathcal{L} (\beta^o) \right\}_i - \varepsilon^2 \sum_{j=1}^{K_{\beta}} \left\{ G^{-1} (\beta^o) \frac{\partial G (\beta^o)}{\partial j} G^{-1} (\beta^o) \right\}_{ij} + \frac{\varepsilon^2}{2} \sum_{j=1}^{K_{\beta}} \left\{ G^{-1} (\beta^o) \frac{\partial G (\beta^o)}{\partial j} \right\}_{ij} \text{tr} \left\{ G^{-1} (\beta^o) \frac{\partial G (\beta^o)}{\partial j} \right\} \]
\[ + \left\{ \varepsilon \sqrt{G^{-1} (\beta^o)} \xi_{i} \right\}, \]
where \( \beta^o \) is the current draw. The proposal density is
\[ q (\tilde{\beta} | \beta^o) = N_{K_{\beta}} (\tilde{\beta}, \varepsilon^2 G^{-1} (\beta^o)) \]
and convergence to the invariant distribution is ensured by using the standard form Metropolis-Hastings probability
\[ \min \left\{ 1, \frac{p (\tilde{\beta} \mid \beta^o, \mathcal{Y}) q (\beta^o | \tilde{\beta})}{p (\beta^o \mid \tilde{\beta}, \mathcal{Y}) q (\tilde{\beta} | \beta^o)} \right\} \]

Again, we can easily generalize this subsection to handle bad inputs and bad outputs. Simply return to the beginning of B.2.1 and include the appropriate price equations for bad inputs and bad outputs, equations (51) and (52), and proceed to generalize the notation.
C.2.2: Drawing Directions

The directions appear in the system in (49) and (50) along with the proposed priors in (39), (C.1.5), and (C.1.6). Using $p_{n,it}$ and $p_{m,it}$ as the left-hand-side variables (which, depending on the system we estimate, may be exogenous or endogenous) we can write (49) and (50) as:

\[
\begin{pmatrix}
  p_{n,it} \\
  p_{m,it}
\end{pmatrix} = \begin{bmatrix}
  -(p_{n,it} + \xi_{it}') & (p_{m,it} + \xi_{it}') \\
  -(p_{n,it} + \xi_{it}') & -(p_{m,it} + \xi_{it}')
\end{bmatrix} \begin{bmatrix}
  \frac{\partial D_{n}}{\partial x_{n}} \\
  \frac{\partial D_{m}}{\partial y_{m}}
\end{bmatrix} \begin{bmatrix}
  g_{n,it} \\
  g_{m,it}
\end{bmatrix} - \begin{bmatrix}
  \xi_{it}' \\
  \xi_{it}'
\end{bmatrix} + \begin{bmatrix}
  v_{n,it} \\
  v_{m,it}
\end{bmatrix}
\]

or, in more compact notation,

\[
p_{it} + \xi_{it} = Q_{it}g_{i} + v_{it}
\]

where $Q_{it}$ is the product of the first two matrices on the right-hand side and $v_{it} \sim N(0, \Sigma_{*})$, where $\Sigma_{*}$ is the relevant submatrix of $\Sigma$ corresponding to $v_{it}$ from the stochastic specification in (C.1.3). From (C.2.2.2), (39), (C.1.5), and (C.1.6) we can draw the directions $g_{i}$ sequentially from the posterior conditional distribution

\[
p(g_{i} | ..., Y) \propto \exp \left[ -\frac{1}{2} (g_{i} - \hat{g}_{i})' V^{-1} (g_{i} - \hat{g}_{i}) \right] \prod_{t=1}^{T} J (g_{i}, \xi_{it}, \gamma)
\]

where $J (g_{i}, \xi_{it}, \gamma)$ is the relevant term of the Jacobian from (C.1.6). The first term comes from (C.1.5) and the next to last term comes from (39).

The first three terms of the conditional posterior can be combined to construct a normal distribution for each $g_{i}$. The Jacobian term can be accommodated either (a) using a Metropolis-Hastings update or (b) using a GC update. Completing the square using the first three terms we obtain:

\[
p(g_{i} | ..., Y) \propto \exp \left[ -\frac{1}{2} (g_{i} - \bar{g}_{i})' V^{-1} (g_{i} - \bar{g}_{i}) \right] \prod_{t=1}^{T} J (g_{i}, \xi_{it}, \gamma)
\]

where

\[
\bar{g}_{i} = \left[ \frac{1}{\Sigma_{*}} A (\gamma)' A (\gamma) + \Psi' \Sigma_{*}^{-1} Q + \Sigma_{g}^{-1} \right]^{-1} (\Psi' \Sigma_{*}^{-1} p_{i} + \Sigma_{g}^{-1} \bar{g})
\]

\[
V = \left[ \frac{1}{\Sigma_{*}} A (\gamma)' A (\gamma) + \Psi' \Sigma_{*}^{-1} Q + \Sigma_{g}^{-1} \right]^{-1}
\]

where $p_{i} = [p_{it} + \xi_{it}, t = 1, ..., T]'$, $\Psi = \text{diag} (Q_{i1}, ..., Q_{iT})$, and $Q_{it}$ has been defined.
The value $c = 10^{-4}$ is fixed so that the technology DDF constraints hold “exactly” (in a numerical sense). Trying lower values of $c$ did not produce any differences in the results that we have obtained.

Once again, we can easily include bad inputs and bad outputs. Return to the beginning of this subsection, add (51) and (52), and generalize the notation.

C.2.3: Drawing Price Distortions

Drawing price distortions $\xi_{it}$ follows essentially the same principles. Assuming that we omit bad inputs and bad outputs, we write the system consisting of (49) and (50) in the following form:

$$
\begin{bmatrix}
    p_{it}^n \\
    p_{it}^m
\end{bmatrix}
= \pi (p_{it}, g_{i})^{t_{N+M}} + \begin{bmatrix}
    -g'_{n,i} & g'_{m,i} \\
    g'_{n,i} & -g'_{m,i}
\end{bmatrix} \begin{bmatrix}
    \frac{\partial D_{it}}{\partial x_n} & 0 \\
    0 & \frac{\partial D_{it}}{\partial y_m}
\end{bmatrix} \begin{bmatrix}
    \xi_{it}^n \\
    \xi_{it}^m
\end{bmatrix} + \begin{bmatrix}
    v_{it}^n \\
    v_{it}^m
\end{bmatrix}
$$

or more compactly as

$$
\mathbf{f}_{it} = p_{it} - \pi (p_{it}, g_{i})^{t_{N+M}} = B_{it} \xi_{it} + v_{o,it} \tag{C.2.3.2}
$$

where $t_{N+M}$ is a $(N + M \times 1)$ unit vector and $v_{o,it}$ is the second block of elements in (C.1.3) that is $v_{it} = \begin{bmatrix} v_{it}, v'_{o,it} \end{bmatrix}$ and its covariance is the lower $(N + M) \times (N + M)$ part of $\Sigma$ defined in (C.1.3), which we denote $\Sigma_o$. We define $B_{it}$ as the product of the two square matrices minus $t_{N+M}$. Given this notation from the joint posterior in (C.1.6) we have:

$$
p (\xi_{it}, \ldots, \mathbf{y}) \propto \exp \left[ -\frac{1}{2} \sum_{i,t} (\mathbf{f}_{it} - B_{it} \xi_{it})^{\prime} \Sigma^{-1}_o (\mathbf{f}_{it} - B_{it} \xi_{it}) - \frac{1}{2} \xi_{it}^{\prime} \Omega^{-1} \xi_{it} \right] \mathcal{J} (g_i, \xi_{it}, \gamma) \tag{C.2.3.3}
$$

The terms in the exponential can be combined, using completion of squares, to provide

$$
p (\xi, \ldots, \mathbf{y}) \propto \exp \left[ -\frac{1}{2} (\xi - \hat{\xi})^{\prime} V_{\xi}^{-1} (\xi - \hat{\xi}) \right] \prod_{i,t} \mathcal{J} (g_i, \xi_{it}, \gamma) \tag{C.2.3.4}
$$

where

$$
\hat{\xi} = \left[ B^{\prime} (\Sigma_o^{-1} \otimes I) B + \Omega^{-1} \right]^{-1} B^{\prime} (\Sigma_o^{-1} \otimes I) \mathbf{f}, \quad \text{and} \quad V_{\xi} = \left[ B^{\prime} (\Sigma_o^{-1} \otimes I) B + \Omega^{-1} \right]^{-1}.
$$

Fortunately, the vector of distortions can be drawn as a block from the above multivariate normal proposal but in order to maintain the correct posterior we have to account for the Jacobian term. Again this can be done using (a) A Metropolis-Hastings update.
or (b) a GC update.

Given a sample \(\{\xi^{(s)}, s = 1, \ldots, S\}\) from the posterior final estimates of price distortions are computed as

\[
\hat{\xi} = S^{-1} \sum_{s=1}^{S} \xi^{(s)}.
\] (C.2.3.5)

Assuming that one wishes to include bad inputs and bad outputs, generalize the notation of this subsection in the same manner as for the previous subsections.

C.2.4: Drawing \(\pi\) and \(\Delta\)

From the joint posterior we have:

\[
p(\pi|\ldots, \mathcal{Y}) \propto \exp\left[-\frac{1}{2} \sum_{i=1}^{N} \sum_{t=1}^{T} (\tilde{z}_{it} - W_{it}^{*}\pi)^{\prime} \Delta^{-1} (\tilde{z}_{it} - W_{it}^{*}\pi)\right]
\] (C.2.4.1)

which is in the form of a multivariate normal. The posterior conditional distribution of \(\Delta\) has the same form as the posterior conditional of \(\Sigma\) and \(\Omega\), that we have described before. One important issue concerns the choice of Fourier terms \((F\) is set to 5). In preliminary experiments we chose \(F\) using the Schwarz information criterion. An alternative approach would have been to consider the Bayes factors for different values of \(F\). Some preliminary investigation revealed that the choice based on the Bayesian Information Criteria (BIC) was fairly robust.

Using the notation in (B.5) and (C.2.4.1) we have:

\[
\pi|\ldots, \mathcal{Y} \sim N_{K+1}(\hat{\pi}, \hat{V})
\] (C.2.4.2)

where

\[
\hat{\pi} = [W_{s}^{\prime} (\Delta^{-1} \otimes I) W_{s} + \overline{V}^{-1}]^{-1} [W_{s}^{\prime} (\Delta^{-1} \otimes I) \tilde{z}]
\] (C.2.4.3)

and

\[
\hat{V} = [W_{s}^{\prime} (\Delta^{-1} \otimes I) W_{s} + \overline{V}^{-1}]^{-1}
\] (C.2.4.4)

where \(\overline{V}\) is the prior covariance matrix of the reduced form parameters defined in (C.2.4.1) and (B.7).

C.2.5: Drawing Technical Inefficiency

We draw \(\{u_{it}\}\) following Tsionas (2006). A technical improvement on that procedure is that we avoid conditionings of the form \(u_{it}|u_{i,t-1}, u_{i,t+1}, \ldots\) and we use another
application of GC where, however, the proposal is the same as in
\[ u_{it}|..., \mathcal{Y} \sim \mathcal{N}^+ (m_{it}, \sigma^2_{it}) \]  
(C.2.5.1)

where \( m_{it} = -\frac{\sigma_{11}\sigma^2_{it}}{\sigma_{11} + \sigma^2_{it}} X^t_{1, it} \gamma, \sigma^2_{it} = \frac{\sigma_{11}\sigma^2_{it}}{\sigma_{11} + \sigma^2_{it}} \). \( X^t_{1, it} \) denotes the right hand side variables of the DDF, and \( \sigma_{11} \) denotes the scalar upper-left element of \( \Sigma \). Drawing \( \gamma_1, \ldots, \gamma_5 \) and \( \gamma_0 \) in (31) is simple and involves only standard results for the normal linear model.

C.2.6 Drawing \( \omega_{it} \)

So far we proceeded under the standard assumption that \( \omega_{it} = 0 \), an assumption that is often if not always used in stochastic frontier analysis and the analysis of distance functions. To generate productivity we take into account (30) with the following modification in (C.1.6), the joint posterior distribution:

- redefine \( V_{it} = \begin{bmatrix} v_{it,1} + \omega_{it} \\ v_{it,2} \\ \vdots \\ v_{it,N+M} \end{bmatrix} = \bar{V}_{it}^* + \omega_{it} \mathbf{1}_{N+M}, \) where \( \bar{V}_{it}^* \) is the old definition of \( V_{it} \) without \( \omega \).

- multiply (C.1.6) by \( \prod_{i=1}^n \prod_{t=1}^T p(\omega_{it} | \omega_{i, t-1}, \bar{p}_{it}^*, \pi_{it}, \tau_{it}, t) \)

- add \( \omega \) to the first element of the last array of equations (C.2.1.2) and (C.2.1.3).

This results from the assumption that the error term of \( \omega_{it} \) say \( \varepsilon_{it,1} \sim \mathcal{N}(0, \sigma^2_{\omega}) \) where the prior of \( \sigma_{\omega} \) is IG: \( \frac{0.01}{\sigma^2_{\omega}} \sim \chi^2_1 \).

To avoid needless technical detail we should notice the following: i) The alternative specifications in (32) (a neural network approximation), (33) (a second-order approximation), (34) (a Fourier approximation) and (35) (a full third-order approximation) are complicated functions of \( \omega_{i, t-1} \); ii) All these specifications depend on \( \pi_{it} \) and \( \bar{p}_{it}^* \) which affect the generation of latent prices and also the directions, through (12)–(15). Given (16) we can ignore the dependence of \( \varrho_{it} \) on directions and use \( \varrho_{it} \) as an observed variable.

These observations raise two technical issues:

- a) How to generate \( \omega_{it} \) conditional on all other parameters and latent variables.

- b) What changes are required in the drawing of all other parameters and latent variables conditional on \( \omega_{it} \).

Regarding (b) we keep the draws obtained when \( \omega_{it} = 0 \), an assumption that should not be too far from the truth. Then, the draws are SIR-reweighted using the appropriate terms in (C.1.6). Overall, the acceptance rates are over 90%, indicating that such approximations are accurate for practical purposes. Regarding (a), \( \omega_{it} \) does not appear
in any of the first-order conditions like (12)–(15) or the semi-informative priors on directions in (27). Therefore, the technical problem is somewhat standard, in that we have to draw from a nonlinear dynamic latent variable model as in (30) and its variations in (32), (33), (34), (35). The consensus in the literature is that Hessian-based approximations followed by standard MCMC are quite accurate. Therefore, we proceed using another update through the GC procedure. All first- and second-order derivatives can be computed analytically using (32)–(35). More importantly, the GC procedure draws \( \{\omega_{it}, t = 1, \ldots, T\} \) as a block for each \( i \). Implementation is similar to drawing \( \{u_{it}\} \) under the specification in (31) although in (31) the model is linear and, therefore, considerably simpler. For technical details see McCausland et al. (2011) and McCausland (2012).

In the implementation of GC we have used a variety of methods for computing first- and second-order derivatives. We report below relative numerical efficiency (RNE) and differences in final posterior means of \( \omega_{it} \) which we use for our statistical inferences on productivity growth. From the results it turns out that: i) It is not essential but it is quite helpful to provide analytical second-order derivatives. ii) AGDH (analytical gradient and a diagonal approximate Hessian) does not work efficiently. iii) ACF and RNE behave similarly irrespective of whether we use analytical or finite-difference approximations to first- and second-order derivatives. Although we have not used it, we also report results for a first-order approximation of \( f \) in (30).

The differences in posterior means when AGDH is used, an implementation that is close to Metropolis-Hastings are large and make this technique rather unlikely to yield reliable results in practice. Only for the linear specification of \( f \) in (30), the posterior means are somewhat close; in all other models we observe huge differences.

### C.2.7: Technical Details

The sequence of MCMC draws can be summarized as follows:

1. Draw \( \gamma \) using the procedures in B.2.1.

2. Draw directions \( g_i \) using the technique in B.2.2, specifically (C.2.2.4).

3. Draw price distortions \( \xi_{it} \) using the technique in B.2.3, specifically (C.2.3.4).

4. Draw parameters \( \pi \) and \( \Delta \) using the method in B.2.4, specifically (C.2.4.1) and (C.2.4.2).

5. Draw technical inefficiency \( u_{it} \) using the method in section B.2.5.

6. Draw covariance matrices \( \Sigma \) and \( \Omega \) using (C.1.6) and (C.1.7).

All Metropolis-Hastings updates use the covariance matrix of the (multivariate normal (MN)) proposal scaled by a constant \( h \) which is selected during the course of the transient or burn-in phase to maintain an overall acceptance rate of 25% for all blocks.
that need Metropolis-Hastings updates due to the complicated Jacobian term. This is done for parameters $\gamma, \xi, g$. For these blocks we compare the numerical performance of the multivariate normal with that of the GC update. The results are summarized in the following Table.

### Table C1.

<table>
<thead>
<tr>
<th></th>
<th>MN for $\gamma$</th>
<th>GC for $\gamma$</th>
<th>MN for $\xi$</th>
<th>GC for $\xi$</th>
<th>MN for $g$</th>
<th>GC for $g$</th>
</tr>
</thead>
<tbody>
<tr>
<td>ACF(50)</td>
<td>0.647</td>
<td>0.325</td>
<td>0.732</td>
<td>0.210</td>
<td>0.725</td>
<td>0.310</td>
</tr>
<tr>
<td>RNE</td>
<td>0.143</td>
<td>0.415</td>
<td>0.225</td>
<td>0.551</td>
<td>0.202</td>
<td>0.562</td>
</tr>
<tr>
<td>$B$</td>
<td>150,000</td>
<td>70,000</td>
<td>150,000</td>
<td>15,000</td>
<td>150,000</td>
<td>7,500</td>
</tr>
<tr>
<td>ACF(50)</td>
<td>0.717</td>
<td>0.344</td>
<td>0.821</td>
<td>0.132</td>
<td>0.781</td>
<td>0.201</td>
</tr>
<tr>
<td>RNE</td>
<td>0.177</td>
<td>0.414</td>
<td>0.124</td>
<td>0.701</td>
<td>0.120</td>
<td>0.661</td>
</tr>
<tr>
<td>$B$</td>
<td>500,000</td>
<td>15,000</td>
<td>400,000</td>
<td>7,500</td>
<td>750,000</td>
<td>9,500</td>
</tr>
<tr>
<td>ACF(50)</td>
<td>0.892</td>
<td>0.332</td>
<td>0.774</td>
<td>0.151</td>
<td>0.816</td>
<td>0.127</td>
</tr>
<tr>
<td>RNE</td>
<td>0.079</td>
<td>0.415</td>
<td>0.137</td>
<td>0.715</td>
<td>0.085</td>
<td>0.664</td>
</tr>
<tr>
<td>$B$</td>
<td>550,000</td>
<td>25,000</td>
<td>450,000</td>
<td>35,000</td>
<td>500,000</td>
<td>15,000</td>
</tr>
</tbody>
</table>

Notes: MN stands for the multivariate normal and GC for the GC update. We discriminate between MN and GC updates for each block individually (see first row). ACF(50) stands for the value of the autocorrelation function at lag 50, RNE for Geweke’s (1992) relative numerical efficiency diagnostic (which should be 1 if i.i.d. sampling from the posterior were feasible) and $B$ denotes the number of MCMC passes required for convergence for the particular block of parameters. In columns reported are ACF(50), RNE, and $B$ for each block of parameters. For each block all statistics reported are medians across all individual elements involved in the block.

From the results it turns out that the numerical behavior of GC updates is impressively better compared to Metropolis-Hastings updates from MN distributions. This fact can be attributed to the substantial dependence of the shape of the posterior on the Jacobian and, of course, its contribution to first- and second-order derivatives of the log-posterior. Reporting of all final results was, therefore, based on the GC updates. Although they converge fast we take a minimum length of the burn-in phase equal to $B=100,000$ followed by another 500,000 MCMC passes which we use to monitor acceptance rates (which remained stable at 20-30%) and compute posterior statistics.

Monotonicity conditions for the technology DDF are checked for every iteration of the MCMC scheme. Draws which result in more than 1% violations are rejected. The average number of rejections per iterations ranged from 500 to 2,500 during the transient phase and from 100 to 500 during the course of MCMC iterations that we finally keep for convergence.