Integrative analysis of gene expression patterns predicts specific modulations of defined cell functions by estrogen and tamoxifen in MCF7 breast cancer cells

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Abstract

To explore the mechanisms whereby estrogen and antiestrogen (tamoxifen (TAM)) can regulate breast cancer cell growth, we investigated gene expression changes in MCF7 cells treated with 17β-estradiol (E2) and/or with 4-OH-TAM. The patterns of differential expression were determined by the ValiGen Gene IDentification (VGID) process, a subtractive hybridization approach combined with microarray validation screening. Their possible biologic consequences were evaluated by integrative data analysis. Over 1000 cDNA inserts were isolated and subsequently cloned, sequenced and analyzed against nucleotide and protein databases (NT/NR/EST) with BLAST software. We revealed that E2 induced differential expression of 279 known and 28 unknown sequences, whereas TAM affected the expression of 286 known and 14 unknown sequences. Integrative data analysis singled out a set of 32 differentially expressed genes apparently involved in broad cellular mechanisms. The presence of E2 modulated the expression patterns of 23 genes involved in anchors and junction remodeling; extracellular matrix (ECM) degradation; cell cycle progression, including G1/S check point and S-phase regulation; and synthesis of genotoxic metabolites. In tumor cells, these four mechanisms are associated with the acquisition of a motile and invasive phenotype. TAM partly reversed the E2-induced differential expression patterns and consequently restored most of the biologic functions deregulated by E2, except the mechanisms associated with cell cycle progression. Furthermore, we found that TAM affects the expression of nine additional genes associated with cytoskeletal remodeling, DNA repair, active estrogen receptor formation and growth factor synthesis, and mitogenic pathways. These modulatory effects of E2 and TAM upon the gene expression patterns identified here could explain some of the mechanisms associated with the acquisition of a more aggressive phenotype by breast cancer cells, such as E2-independent growth and TAM resistance.

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Introduction

Estrogens play several major roles in mammalian physiology, including the control of the reproductive tract and development of secondary sex organs. They regulate the estrous cycle, control lactation in the mammary gland and affect bone, liver and cardiovascular systems (Sutherland et al. 1988, Couse & Korach 1999). In diseases such as breast cancer, estrogens act as mitogenic factors and are important in tumor initiation and progression in vitro and in vivo (Lippman et al. 1975, Soule & McGrath 1980, Key & Pike 1988, Colditz 1998).

Studies on estrogen receptor (ER)-positive breast cancer cell lines indicate that estrogens and antiestrogens act on cell populations in early to mid-G1 phase (Sutherland et al. 1983, Leung & Potter 1987). It is commonly accepted that tamoxifen (TAM) binds to the ER and inhibits ER-mediated gene transcription. Although antiestrogens are widely employed in the treatment of hormone-responsive breast cancer to induce cell growth arrest, the mechanisms by which TAM or its derivatives regulate gene expression are not well understood. Understanding of these mechanisms could allow us to explain, at least in part, the development of cellular resistance to TAM treatment and consequent therapeutic escape (Gibson et al. 1990, Katzenellenbogen 1991, Osborne et al. 1995, El Etreby & Liang 1998).

To explore, at gene expression level, how estrogen and antiestrogen could regulate cell growth and tumor progression in breast cancer, we investigated the...
patterns of gene expression in MCF7 cells treated with TAM and/or with 17β-estradiol (E2). The MCF7 cells, isolated from a pleural effusion of metastatic human breast adenocarcinoma, are ER and progesterone receptor (PR) positive. Their growth is estrogen dependent and sensitive to TAM (Soule et al. 1973, Sommers et al. 1990).

The patterns of differential gene expression associated with E2-induced tumor progression and with TAM antitumor action were analyzed by the ValiGen Gene IDentification (VGID) process. This process is based on a subtractive hybridization method combined with microarray and integrative data analyses that have previously been described (Gadal et al. 2003). The subtractive hybridization method does not require specific probes or primers to isolate differentially expressed mRNAs, thus allowing identification of isoforms and unknown transcripts without having to predetermine the sequences to be investigated. Microarrays provide a rapid and high throughput screening method for validating the differentially expressed status of sequences isolated by the subtractive hybridization process. Differential expression data and extensive published information were then integrated into a detailed biologic model. First, we generated a relational graph encompassing all known functional interactions between genes, proteins and small molecules recorded in the scientific literature pertaining to mammalian cellular and physiologic mechanisms. Then, the entire differential expression data set was injected into the graph in order to extract a subgraph functionally correlating the data sets. This enabled us to construct physiologic/signaling interaction maps directly correlated with gene expression patterns and protein interactions (Fig. 1). For more details, see Gadal et al. 2003 and our web page: http://perso.club-internet.fr/fgadal. The resulting model points out some of the mechanisms through which E2 and TAM could affect the initiation, growth and progression of breast cancer.

Thus, by combining subtractive hybridization, cDNA microarrays and data integration, we were able to highlight some of the interconnected mechanisms, such as cytoskeleton remodeling, matrix degradation, cell cycle progression, synthesis of genotoxic metabolites, DNA repair and estrogen receptor constitution/activity, through which estrogen and antiestrogen regulate tumor progression.

Materials and methods

Cells and cell culture

The MCF7 breast cancer cell line, obtained from ATCC (Rockville, MD, USA), was cultured routinely in Glutamax I Dulbecco’s modified Eagle’s medium (DMEM) (Gibco Invitogen) supplemented with 10% fetal calf serum (FCS) (Gibco Invitogen), 50 µg/ml streptomycin (Gibco Invitogen) and 50 UI/ml penicillin (Gibco Invitogen) at 37 °C in 5% CO2 atmosphere. The experiments were carried out in phenol red-free DMEM (Gibco Invitogen) supplemented with 10% charcoal-treated FCS, 50 µg/ml streptomycin, 50 UI/ml penicillin and 2 mM L-glutamin (Gibco Invitogen). The cells, divided into three batches, were incubated at 37 °C in 5% CO2 atmosphere for 8 days. Then, one batch of...
MCF7 cells was stimulated with $1 \times 10^{-8}$ M E$_2$ (Sigma Aldrich) for 15 h, whereas the second batch was maintained in E$_2$-free medium. To explore E$_2$-associated differential expression, the VGID procedure was performed with cDNAs generated from batches 1 and 2. To explore TAM-associated differential expression, the VGID procedure was performed with cDNAs generated from batches 1 and 2. The third batch of MCF7 cells was incubated with $1 \times 10^{-8}$ M E$_2$ and $1 \times 10^{-7}$ M TAM (Sigma Aldrich) for 15 h, and the VGID procedure was performed with cDNAs generated from batches 2 and 3.

**cDNA synthesis**

Direct mRNA capture from lysed cells was performed with the Dynabeads mRNA direct kit (Dynal, Compiegne, France) according to the manufacturer’s protocol. First-strand cDNA synthesis was carried out as follows: 6 µl aliquots of first-strand synthesis mixture (10 mM dNTP each (Promega), 0-1 M DTT and 5 first-strand buffer (both from Life Technologies, Cergy Pontoise, France)), 500 ng Dynabeads-bound denatured mRNA and 200 units Superscript II (Life Technologies) were separately preincubated at 37°C for 1 min in a water bath, and then mixed and incubated in a final volume of 30 µl at 37°C for 1 h. The first-strand samples were then transferred to 170 µl diluted second-strand synthesis mixture (1 final E. coli DNA ligase buffer (New England Bio-Labs, Saint Quentin Yveline, France), 900 mM KCl (Sigma Aldrich), 20 mg/ml glycogen (Boehringer Mannheim, Mannheim, Germany), 10 mM dNTP each, 10 U/µl E. coli DNA ligase (New England Bio-Labs), 2 U/µl RNase H and 10 U/µl E. coli DNA polymerase I, all purchased from Promega) and incubated for 2 h at 16°C in a PerkinElmer (Courtaboeuf, France) 9700 thermal cycler. The cDNA populations, still bound to Dynabeads, were then digested with Sau3A (4 U/µl; New England Bio-Labs) for 2 h at 37°C to generate the fragments of 256 base pairs in statistical length. The reaction was stopped by heating for 20 min at 65°C. The digested cDNAs were then ligated to adapters, and PCR was performed with adapter-specific primers in a PerkinElmer 9700 thermal cycler, using 12 cycles of 95°C for 30 s, 55°C for 45 s and 72°C for 1 min, followed by 72°C for 7 min. The PCR products were purified by the Qiaquick PCR purification kit (Qiagen) and used for VGID and/or microarray experiments (see below).

**Adapters and oligonucleotides**

Sequences of adapters (CyberGene, Saint Malo, France) used in VGID procedure were as follows: BamHI adapter, 5’ GTT AGA ACG AGA CGG ATC CTG 3’ and 3’ TT GAA TCT TGC TCT GCC TAG GAC TAGp5; BglII-bio adapter, bio 5’CCA GCT AAC ACC TAG ATC TGC 3’ and 3’ TT GGT CGA TTG TTG TGT3’. The fragments of cDNA ligated to these adapters were amplified with the corresponding oligonucleotides: BamHI primer, 5’ AA CTT AGA ACG AGA CGG ATC CTG ATC 3’ and BglII-bio primer, bio 5’ AA CCA GCT AAC ACC TAG ATC TGC ATC 3’. The inserts were amplified with the primers pVG17 (5’ CTT CCG GCT GCT ATG GTA 3’) and pVG17 reverse (5’ CTT CCG GCT GCT ATG GTA 3’).

**VGID gene identification technology (patent no. 6221585)**

As previously described (Gadal et al. 2003), VGID directly isolates overexpressed and underexpressed cDNA associated with the transition from a defined phenotypic state to another state within a congenic system. For the first denaturation-renaturation step, 300 ng BamHI-ligated-amplified tester were mixed with 1200 ng BglII-bio-ligated-amplified driver. The mixture was ethanol-precipitated, resuspended in 4 µl HEPES 0.5 M-EDTA 0.2 mM and overlaid with 20 µl mineral oil, denatured 5 min at 98°C and finally chilled on ice. The salt concentration was adjusted to 0.5 M with 1 µl of 2.5 M NaCl. After 5 min denaturation at 98°C, the sample was allowed to anneal for 20 h at 65°C. After hybridization, the oil was removed. The sample volume was adjusted to 100 µl with 100 mM NaCl and 10 mM Tris HCl, pH 8, and mixed with 210 µl streptavidin magnet beads (10 mg/ml, Boehringer Mannheim) to recover tester single- and double-strand DNA, as described by the manufacturer. This step was repeated to remove all biotinylated cDNA. To recover the tester, double-strand cDNA, the supernatant, which contained unbiotinylated cDNA, was incubated with 1.5 µg single-strand binding protein (SSB, Promega) for 30 min at room temperature. The sample was loaded onto a Millipore Micropure EZ membrane (which retained the proteins) and centrifuged for 1 min at 20 800 g at room temperature. The flow through was combined with 1200 ng driver, and the next round of hybridization was set up as described above. A total of three hybridization rounds were performed. The cDNA recovered after the last round, was amplified by PCR, as described above, using 25 cycles and the BamHI primer. The sequences under- or overexpressed were cloned into a modified pUC19 vector (Gibco Invitrogen).

**Cloning**

The vector used for the cloning step was a derivative of the pUC19 vector in which polylinker EcoRI–HindIII was replaced by the 5’ ATTTCGGATCCTCA3’ pVG17 sequence. To avoid cloning chimeric structures, the vector/insert DNA ratio used was 3:1. Vector (50 ng) was ligated to the recovered cDNA and amplified after

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subtraction in the presence of 1 U ligase (Boehringer Mannheim) in a final volume of 10 µl overnight at 16°C. The ligation mixture was purified by the GeneClean procedure (BIO 101), according to the manufacturer’s recommendation, and the 2 µl-aliquot was used to transform 50 µl MAX Efficiency DH10B cells (Gibco Invitrogen). Transformation was performed by electroporation at 1800 V with the EC100 Electroporator (E-C Apparatus Corporation, Petersburg, FL, USA). Immediately after electroporation, 500 µl SOC medium were added. Cells were shaken for 1 h at 37 °C, and an aliquot was spread on LB medium plate containing 100 µg/ml ampicillin.

**Sequencing**

Isolated colonies were selected in 150 µl LB with 100 µg/ml ampicillin in order to make glycerol stocks. A volume of 4 µl of this solution was cycle-sequenced in a PerkinElmer thermal cycler with the AmpliTaq Gold DNA polymerase kit (Perkin Elmer), using 10 pmol pVG17(-74) and pVG17 rev(-88) primers in a final volume of 50 µl. A pre-PCR step, performed at 95 °C for 13 min, allowed the activation of the polymerase and the bacteria lysis. Each of the 40 PCR cycles included three segments: 95 °C/30 s, 54 °C/1 min and 72 °C/2 min 30 s. The PCR products were sequenced without purification with the BigDye Terminator Cycle Sequencing kit and ABI Prism 3700 automatic DNA sequencer (PerkinElmer Applied Biosystem).

**Sequence analysis**

The clone sequences were analyzed by in-house programs written in Perl. After removal of the vector sequence, the repeats were masked and compared with public databases with the BLASTn and BLASTx programs. The significance of the similarities was checked at both the nucleic and protein level. DNA sequences were considered for further analysis when the level of similarity with known sequences was greater than 98%. At the protein level, significant similarity threshold was fixed as at least 40. The sequence data were then clustered with Fasta software (Infobiogen, Evry, France). Unidentified genes (sequences failing to match anything in the databases, including expressed sequence tags (ESTs)) could not help us to construct the relational graph. Thus, these sequences were not introduced in the analysis.

**Microarray construction**

All the clones obtained from the subtraction process were studied with microarray technology. Each clone was PCR-amplified in a final volume of 100 µl, using 20 pmol of both primers pVG17(-74) and pVG17 rev(-88), 10 nmol dNTPs and 1·5 U DyNazyme EXT. After 13-min enzyme activation at 95 °C, 40 cycles were carried out (95 °C/30 s, 56 °C/1 min and 72 °C/1 min 30 s). A final incubation was performed for 7 min at 72 °C. The PCR products were purified with the Multiscreen PCR Kit (Millipore, Saint Quentin Yveline, France), according to the manufacturer’s instructions, and concentrated under Vacuum SpeedVac (E-C Apparatus). The PCR products, resuspended in 3 SSC, with 11 yeast genes used as internal control, were spotted in triplicate onto GAPS Amino Silane Coated Slides (Corning, Schiphol-Rijk, The Netherlands) with the GMS 417 arrayer (Genetic MicroSystem, Affymetrix, High Wycombe, Herts, UK). The slides were UV cross-linked at 300 mJ, prehybridized in 50% formamide, 0·1% SDS, 1% bovine serum albumin (BSA) and 5 SSC buffer at 42 °C for 1 h, washed first in water and then in 95% ethanol, and finally vacuum dried.

**Microarray hybridization, scanning and data acquisition**

cDNA (500 ng) from MCF7 cells treated or not with TAM or E2 were labeled by random priming with incorporation of Cyanine5-dUTP for the tester DNA and Cyanine3-dUTP for the driver samples respectively. Then, the samples were mixed, concentrated by evaporation under vacuum and resuspended in prehybridization buffer, as described above, with Denhardt’s solution replacing BSA. The two-labeled cDNA mixtures (MCF7 cDNA mixed with MCF7 + E2 one and MCF7 + E2 cDNA mixed with MCF7 + E2 + TAM one) were hybridized with the arrayed slides overnight at 42 °C. The slides were then washed for 5 min with 1 SSC-0·1% SDS, 3 min with 1 SSC, 3 min with 0·1 SSC and 1 min with water, and finally 95% ethanol-dried and scanned (GenePix 4000A; AXON, Union City, CA, USA). Accurate differential measurements (final fluorescence ratios) were expressed as the average of nine independent assays where each sequence was arrayed in triplicate. Visualization, quantification and gene expression analysis were performed with GENEPIX 3·0 software (AXON). The data were normalized by the autonormalization method of Yang et al. (2000).

**Assay of NAD(P)H quinone oxidoreductase and NADH-menadione oxidoreductase activity**

MCF7 cells were cultured, as described above, with E2. The MCF7 cells were harvested with a rubber policeman, centrifuged at 1000 g for 5 min at 4 °C and homogenized in cold buffer (100 mM potassium phosphate, pH 7·0, and 2 mM EDTA). After centrifugation at 10 000 g for 15 min at 4 °C, the supernatant containing cytosolic and microsomal enzymes was isolated. NAD(P)H quinone oxidoreductase activity was

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determined with 2,4-dichlorophenol indophenol (DCPIP) (Sigma). Samples (10 µl) were mixed with 80 µl of 1 mM NAD(P)H (Sigma Aldrich) in a microtiter plate and incubated for 15 min at 37 °C. Then, 4 mM DCPIP (10 µl) was added to each well, and the absorbance was measured at 600 nm every 5 min. To assess NADH-menadione oxidoreductase activity, the samples (10 µl) were mixed with 70 µl of 10 µM menadione (Sigma Aldrich) and 10 µl of 40 µM cytochrome C (Sigma Aldrich). After determining the baseline at 550 nm, 10 µl of 0·5 mM NADH (Sigma Aldrich) was added, and the absorbance was measured every minute. Changes in absorbance were obtained by plotting the absorbance values as a function of time to determine the slope (rate) of the linear portion of the curve. Then, changes in absorbance corresponding to the activity of the enzymes were expressed as percentages.

**Results**

**Differential gene expression patterns in MCF7 cells treated with E₂ or E₂ and TAM**

To determine the sequences overexpressed in response to E₂ in MCF7 cells, the sequences contained in the E₂-treated MCF7 cDNA library (tester) were subjected to competitive hybridization against an excess amount of cDNA from untreated MCF7 cells (driver). This was followed by selective trapping of all driver material, including sequences common to both tester and driver. In order to obtain E₂-underexpressed sequences, the cDNA obtained from E₂-treated MCF7 cells was ligated to BglII-bio adapter to become the driver library, whereas the cDNA from untreated MCF7 cells was ligated to BamHI adapter to form the tester library. Competitive hybridizations were then performed.

In the TAM experiments, the E₂ + TAM-treated MCF7 cDNA library (tester) was subjected to competitive hybridization against an excess amount of cDNA from E₂-treated MCF7 cells (driver). Sequences underexpressed in response to TAM treatment were obtained by inverting the adapter-dependent tester:driver identification tagging, as described above.

All competitive hybridizations were performed with a driver/tester ratio of 4, which allowed optimal recovery of sequences differentially expressed (data not shown). Three iterative rounds of subtraction and selective trapping resulted in high enrichment and efficient isolation of tester-specific sequences, as visualized by migration on agarose gel (Fig. 2). The electrophoretic profiles were composed of smears, ranging from 100 to 1250 pb, as well as characteristic bands corresponding to cDNAs underrepresented (lanes 1 and 3) and overrepresented (lanes 2 and 4) in MCF7 cells treated with E₂ alone (lanes 1 and 2) or in combination with TAM (lanes 3 and 4). E₂ treatment resulted in the identification of three different major cDNAs bands, observed in both lane 1 (underrepresented cDNAs) and lane 2 (overrepresented cDNAs) while treatment with E₂ + TAM led to the identification of nine differentially expressed sequences, five of which were underexpressed (lane 3).

All these differentially expressed sequences were characterized by distinct lengths, suggesting that they reflect different gene expression patterns, which could be specifically associated with differences in the physiologic states of the cells after estrogen or antiestrogen treatments.

All the cDNAs recovered after subtractive hybridization were then cloned in a modified Puc vector. To avoid preferential cloning of smaller cDNAs over larger fragments, the VGID outputs were electrophoretically separated on agarose gels, which were then divided into four parts to separate the cDNAs into size populations. The first group corresponded to cDNAs larger than 700 pb. The sizes of the second and third cDNA groups were 500–700 pb and 300–500 pb respectively. The last section was composed of cDNA fragments less than 300 pb in length. Overall, the subtractive hybridization processes yielded 1005 cDNA inserts, ranging in size from 100 to 1250 bp. Sequence analysis (Table 1) revealed that treatment with E₂ resulted in the differential expression of 279 known and 28 unknown sequences, whereas treatment with E₂ + TAM resulted in the differential expression of 286 known and 14 unknown sequences. Redundant identification of known sequences was relatively low, corresponding to 28–40% of the datasets, depending upon the VGID experiments (Table 1). This strategy allowed us to identify a wide panel of genes involved in different cell functions affected by E₂ and TAM treatments. Here, redundant identification of sequencesgenes did indicate relative expression level, which could then be validated by microarray analysis. For example (Table 2), the pS2 sequence was represented by 50 clones, suggesting substantial overexpression of the gene, whereas the upstream factor 2 (USF2) sequence was found only once, arguing for moderate to low overexpression. Indeed, microarray analysis showed that expression of pS2 and USF2 was enhanced by 4·93- and 1·44-fold respectively.

To confirm the differential expression status of the sequences isolated by VGID and obtain quantitative information, the cDNA insert of all the clones, known as well as unknown nonredundant sequences (607 sequences), were PCR amplified and arrayed on slides. Because of the existing overlap between the two data sets, the 147 genes subject to differential regulation in response to both E₂ and TAM were represented twice on the same microarray. The microarrays thus constructed, representing a total of 460 different probes, were then independently hybridized to each of the different mRNA populations corresponding to our experimental conditions. Thus, when verifying the
differential responses obtained after estrogen treatment, for instance, the microarrays utilized contained not only all the genes identified as estrogen responsive after VGID analysis, but also all those identified as TAM responsive. Multiple randomly selected cDNAs were also spotted on the same slides to serve as internal controls. Post-hybridization analyses were based on the autonormalization method of Yang et al. (2000). For example, the cDNA obtained from E2-treated MCF7 cells was labeled with Cy-5 fluorochrome (red); that from untreated MCF7 cells with Cy-3 fluorochrome (green). After hybridization (Fig. 2), red and green fluorescence indicated greater relative expression in the presence and in the absence of E2 respectively. Yellow fluorescence corresponded to equivalent expression levels. Microarray analyses confirmed both the patterns and levels of differential expression of the majority (~90%) of the genes identified as differentially expressed through

Table 1 Distribution of the MCF7 differentially expressed sequences to response in E2 and TAM treatments. cDNA transcripts obtained by subtractive hybridizations were cloned into a modified Puc vector and sequenced. Gene identification was performed with BLAST software against nucleotide and protein banks.

<table>
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<th>VGID subtractive hybridization</th>
<th>Under-expressed with E2</th>
<th>Over-expressed with E2</th>
<th>Under-expressed with TAM</th>
<th>Over-expressed with TAM</th>
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<tr>
<td>Number of clones sequenced</td>
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<td>236</td>
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<td>Number of known sequences</td>
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<td>207 (148 non-redundant)</td>
<td>221 (138 non-redundant)</td>
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<td>Number of unknown sequences</td>
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<td>8 (8 non-redundant)</td>
<td>7 (5 non-redundant)</td>
<td>9 (non-redundant)</td>
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</table>

Figure 2 Isolation of transcripts differentially expressed in MCF7 cells. Gel migration analyses (1.5% agarose) were performed at each step of the VGID process. Lanes 2 and 4 correspond respectively to cDNA samples of sequences overrepresented in the MCF7 + E2 and MCF7 + E2 + TAM libraries. For identification of the transcripts underexpressed in MCF7 + E2 and MCF7 + E2 + TAM, the process was repeated, using the MCF7 and the MCF7 + E2 cDNA libraries as tester respectively. Thus, lanes 1 and 3 represent sequences underrepresented in the MCF7 + E2 and the MCF7 + E2 + TAM cDNA libraries after differential trapping.
Table 2 Function and expression level of E2-inducible genes. The levels of expression indicate over-expression (upward arrows) and under-expression (downward arrows). These expression patterns were qualitatively and quantitatively verified by microarray expression analyses. Relative expression levels are indicated by arrows and x-fold. The last column indicates the genes whose expression is reversed by TAM treatment in MCF7 cells. In the 'TAM inducible gene expression switching' column, "-" indicates that no switching effects were experimentally observed. Data for all VGID-identified genes can be accessed at the following website address: http://perso.club-internet.fr/fgadal

<table>
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<tr>
<th>Broad functions</th>
<th>E2-responsive genes</th>
<th>Accession number</th>
<th>VGID identification status</th>
<th>Number of clone corresponding to identified genes</th>
<th>Microarray status (X fold expressed)</th>
<th>TAM-inducible gene expression switching</th>
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<td>Cytoskeletal remodelling</td>
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<td>Cell-cycle progression</td>
<td>NAD(P)H menadione oxidoreductase</td>
<td>J03934</td>
<td>/</td>
<td>2</td>
<td>1.30±0.05</td>
<td>1.28±0.01</td>
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<td>UDP-glucuronosyltransferase</td>
<td>M57951</td>
<td>/</td>
<td>1</td>
<td>1.20±0.10</td>
<td>-</td>
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<tr>
<td></td>
<td>Thiol-specific antioxidant</td>
<td>Z22548</td>
<td>/</td>
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<td>1.56±0.02</td>
<td>1.18±0.04</td>
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<tr>
<td></td>
<td>NADH dehydrogenase ubiquinone</td>
<td>NM_005006</td>
<td>/</td>
<td>4</td>
<td>1.91±0.09</td>
<td>1.33±0.01</td>
</tr>
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</table>
utilization of VGID, except for immunophilin (Table 3) (entire data set available at http://perso.club-internet.fr/fgadal).

Integrative analysis of the information embodied in the literature, combined with the VGID and microarray results, singled out a set of 32 differentially expressed genes coding for proteins associated with broad cellular functions (Tables 2 and 3). The expression status (under- or overexpressed), as indicated by the VGID outputs for 31 of these identified genes, was corroborated by microarray analysis. In our dataset, eight genes were found to be differentially expressed by a factor superior to twofold, nine by a factor of 1·50–2·00 and 14 by a factor inferior to 1·50. These results indicate that the VGID procedure is sensitive enough to isolate transcripts the expression level of which is only slightly modified.

E2 modulates biologic functions in MCF7

Integrative analysis indicated that, in MCF7 cells, E2 treatment seemed to affect (Table 2):

(i) Anchors and junctions remodeling. Seven of the eight genes identified as playing significant roles in these mechanisms were repressed, including β catenin, junction plakoglobin, γ-actin-1, claudin 4, the Arp2/3 complex involved in adherent junctions, and keratins 8 and 18, implicated with junction plakoglobin in desmosomes constitution. Only Hsp27, which plays a role in adherent junctions, was found to be upregulated.

(ii) Extracellular matrix (ECM) degradation. In ECM degradation, underexpression of hepatocyte growth factor activator inhibitor 2 (HAI-2) and USF2, together with overexpression of cathepsin D and pS2, argued for an important role of E2 in the activation of proinvasive mechanisms through matrix remodeling.

(iii) Cell-cycle progression. Overexpressed c-myc, nm 23-H2 nucleotide diphosphate kinase, cyclin kinase subunit 1 (Cks1), CDK2, B-myb, activator of S phase kinase (ASK) and underexpressed TGFβ are involved in cell-cycle progression, in particular with the G1/S checkpoint and in S-phase progression (Fig. 3).

(iv) Synthesis of genotoxic metabolites. Genes involved in detoxification mechanisms, such as UDP glucurono-syltransferase, NAD(P)H menadione oxidoreductase and thiol-specific antioxidant, were found to be underexpressed, whereas NAD(P)H dehydrogenase ubiquinone was overexpressed in MCF7 cells after E2 treatment, suggesting a decrease in the efficiency of at least one important detoxification pathway.

E2 decreases the activity of both NAD(P)H menadione oxidoreductase and NAD(P)H quinone oxidoreductase

To verify our conclusion that E2-treatment induced modifications in the activities of the oxidant-dependent and genotoxic quinone-dependent pathways (Fig. 4), we measured the activities of two key enzymes, NAD(P)H menadione oxidoreductase and NAD(P)H quinone oxidoreductase. In E2-treated cells, we observed that activities of both enzymes were reduced by 11±2% and 30±6% (P<0·05) respectively, as compared with untreated MCF7 cells.

TAM antagonizes only some effects of E2 upon gene expression in MCF7 cells

Our integrative data analysis indicated that TAM treatment reverses E2-induced changes in the expression of only 11 genes and apparently restores only some of the biologic functions affected by E2 (Table 2).
Increases in the expression of claudin 4, γ-actin-1, keratin 8, keratin 18 and decrease in the expression of Hsp27 suggest that TAM treatment could restore anchoring and junction structures.

Switching in HGFA I2, cathepsin D and pS2 expression suggests that TAM could restore protection of ECM integrity.

Expression switching of NAD(P)H menadione oxidoreductase, thiol-specific antioxidant, and NADH dehydrogenase ubiquinone argues for a possible restoration of detoxification mechanisms, including a decrease in E2-induced quinones synthesis, which could prevent catechols/quinone genotoxic effects.

Interestingly, we did not find any change in the expression of genes involved in cell-cycle progression previously affected by E2. Furthermore, we found (Table 3) that TAM treatment specifically decreases the expression of the DDB-2 and XR-CC1 genes involved in DNA repair. TAM treatment also inhibits the expression of Hsp90 and immunophilin while increasing that of REA. These three gene products are known to play a role in the formation of active ER. Concurrently, TAM treatment induces the overexpression of profilin, which is associated with actin dynamics. Finally, TAM treatment induces underexpression of TGFβ and overexpression of ERK1 and hydroxysteroid (17-beta) dehydrogenase 4 (HSD17B4).

**Discussion**

In the present study, we used the analytic approach we had previously described (Gadal et al.) to identify the genes differentially expressed in MCF7 breast cancer cells stimulated with E2 or treated with TAM. With this approach, the capture of biotinylated cDNA and single-strand cDNA is very effective, thereby avoiding the construction of normalized cDNA libraries and increasing significantly the number of differentially regulated genes correctly identified. The differential expression pattern of over 90% of the sequences thus isolated was confirmed by microarray analyses. These
techniques have been significantly improved over the past 4 years, and it is now widely accepted that they can reliably be used to verify accurately and independently the relevance of differential expression data generated by new technologies, such as the VGID approach, for instance. For comparison, with Suppression Subtractive Hybridization (SSH) as the primary means to detect differential expression, only 70% of identified genes were confirmed by a microarrays approach similar to that described above (Kuang et al. 1998).

Furthermore, and in opposition to high-density microarrays, the VGID strategy avoids the introduction of analytic bias resulting from the a priori selection of genes to be investigated. We then carried out computer-assisted data integration to achieve an understanding of the physiologic mechanisms potentially affected by differential expression of the identified genes. Initially, the sets of genes identified as estrogen or TAM responsive were completely separate because they were independently obtained from different RNA populations. Subsequent data analysis showed that some genes were differentially modulated under each of our two experimental conditions, thereby indicating partial overlap between the two datasets. The VGID data sets included genes previously shown to be regulated in response to E2 or TAM, such as Hsp27 (Porter et al. 2001), pS2 (Inadera et al. 2000), cathepsin D (Inadera et al. 2000, Safe 2001), c-myc (Doisneau-Sixou et al. 2003), ERK1 (Rabenoelina et al. 2002) and β-catenin (Gunin et al. 2003), thereby substantiating the validity of our experimental approach. The fact that only immunophilin expression changes were not corroborated by microarray analyses may be explained by the observation that quantitative information obtained with small cDNA inserts as well as clones arising from less abundant mRNA species could be unreliable due to their propensity to generate weak signals falling within the ‘noise’ of microarray hybridization signals (Yang et al. 1999). Here, the transcript for

![Figure 4](https://www.endocrinology-journals.org/34,61-75) Evidence of E2 genotoxic effects in MCF7 cells. Boxes with plain color represent genes found to be differentially expressed by the VGID process and confirmed by cDNA microarray analysis.
immunophilin could well belong to the low-abundance class since it was represented by only one isolated clone (Table 3). Despite this limitation, our approach allowed us to isolate and screen a large number of transcripts, pointing out physiologic mechanisms apparently affected by estrogenic or antiestrogenic treatments in MCF7 breast cancer cells.

MCF7 cells develop into xenografts in ovariectomized athymic mice only when supplemented with E2 (Soule & McGrath 1980), indicating that E2 promotes primary tumor growth. Our results show (Table 2) that, in MCF7 cells, E2 regulates the expression of seven genes involved in the transition from the G1 to the S phase of the cell cycle, such as c-myc (Safe 2001, Doisneau-Sixou et al. 2003), CDK2 (Safe 2001, Doisneau-Sixou et al. 2003), b-myb (Saville & Watson 1998), ASK (Masai et al. 2000), CKS1 (Spruck et al. 2001), TGFβ (Polyak et al. 1994) and nm23-H2 (Ji et al. 1995). Five of these, CDK2, nm23-H2, CKS1, b-myb and ASK, are reported here, for the first time, to our knowledge, as being regulated by E2 at mRNA level. CDK2 forms complexes with cyclins E and A, indispensable for progression through mid- and late G1 phase and entry into S phase (Safe 2001) (Fig. 3). Thus, E2-induced increase in CDK2 expression should enhance the level of CDK2 protein available to form these complexes. The cyclin A/CDK2 complex was reported to activate b-myb involved in S-phase progression (Saville & Watson 1998). Overexpression of b-myb, observed here, could reinforce cell-cycle progression. Interestingly, we also observed the overexpression of ASK, another gene implicated in cell-cycle progression. Interestingly, we also observed the overexpression of ASK, another gene implicated in S-phase progression (Masai et al. 2000). Here, CKS1 induces the degradation of p27, a CDK2 inhibitor (Spruck et al. 2001), thus increasing CDK2 activity. The overexpression of CKS1 can also result from TGFβ underexpression, since TGFβ inhibits CKS1 transcription (Simon et al. 1995). Finally, decrease in TGFβ level induces additional inhibition of p27 expression (Polyak et al. 1994, Reynisdottir et al. 1995). Altogether, these different effects converge to enhance CDK2 activity in addition to the increases in CDK2 mRNA levels described above. In this study, we also observed the E2-induced overexpression of nm23-H2, a nucleotide diphosphate kinase activating c-myc expression (Ji et al. 1995). This suggests that E2 directly and indirectly stimulates c-myc expression.

One of the key features of an aggressive phenotype is the ability of the cancer cells to escape from the tumor stroma, thus promoting metastasis. The invasive process through the basement membrane requires changes in intercellular adhesion, cell motility and remodeling of the ECM (DePasquale et al. 1994, Lochter & Bissell 1995). In epithelium, desmosomes form strong adhesive junctions between cells by linking intermediate keratin filament networks to sites of intercellular adhesion. An essential structural component of desmosome is junction plakoglobin, the underexpression of which is associated with anchorage loss in cancers (Tada et al. 2000). Our results show that E2 inhibits the expression of junction plakoglobin as well as that of keratins 8 and 18, which are interconnected to desmosomes via binding to desmoplakin, arguing for E2-induced desmosome disorganization. In agreement with this, DePasquale et al. (1994) have observed a rearranging of vesicular plakoglobin staining after E2 treatment of MCF7 cells, which could be a consequence of plakoglobin underexpression. We also observed in E2-treated MCF7 cells, for the first time, to our knowledge, the underexpression of subunit 1B of the Arp2/3 complex, which is thought to promote actin filament assembly at the barbed end (Cooper & Schafer 2000), suggesting E2-induced retardation of actin microfilament polymerization. Furthermore, the overexpression of Hsp27 (Porter et al. 2001) (Table 2), which inhibits the formation of basolateral microfilaments, suggests that E2 treatment could alter microfilament homeostasis and microfilament dynamics (Piotrowicz & Levin 1997). Catenins form an important complex with cadherins, linking them to the actin filament network. Underexpression of β catenin (Gunin et al. 2003) in E2-treated MCF7 cells could modify this process. Indeed, decrease in cadherin-associated β catenin has been associated with invasiveness in breast cancer (Gonzalez et al. 1999). Importantly, E2 decreased γ-actin expression (Table 2), a phenomenon known to be associated with the progression from a nonmetastatic to a metastatic phenotype of cells in salivary gland adenocarcinoma (Suzuki et al. 1998). Finally, we observed, for the first time, to our knowledge, E2-regulated expression of claudin 4, involved in tight junction constitution, the underexpression of which is associated with increased invasive potential (Michl et al. 2003). Altogether, our results and those of other workers indicate that, in MCF7 cells, E2 treatment modulates the expression of major genes promoting the remodeling of cell adhesion structures, and leads, in consequence, to decreases in cell adhesiveness and increases in cell mobility, thereby setting off two major causal mechanisms associated with tumor progression and the acquisition of an invasive potential (Suzuki et al. 1998) through the loss of stable cell–cell adherent junctions (DePasquale et al. 1994).

In this study, we identified E2-induced overexpression of cathepsin D and pS2, and underexpression of two new genes, USF2 and HGFA II. Known to be associated with invasive human breast cancer (Elliott et al. 2002) and metastasis (Birchmeier et al. 1997), HGF can activate the transcription factor E1AF (Hanzawa et al. 2000), which, in turn, stimulates the production of various MMPs (matrix metalloproteinases) (Hanzawa et al. 2000), thereby promoting matrix degradation. HGF is synthesized as pro-HGF/SF, and it requires activation by factors such as HGF activator (HGFA).
Thus, through a decrease in HGFA L2 (inhibitor of HGFA) expression, E2 appears to promote HGFA-mediated specific activation of HGF, resulting in ECM degradation. This process could be reinforced by the underexpression of USF2, observed here for the first time, which interacts, like other USF transcription factors, with the E1 box binding site on the HGF gene multiconensus region. Jiang et al. (2000) have shown that only USF-1, and not USF-2, can stimulate HGF transcription. We can speculate that underexpression of USF2 could diminish USF-2 competition for E1 box, facilitate USF-1-dependent HGF gene transcription and promote E1AF-dependent MMPs gene transcription, leading to increased matrix degradation potential.

Furthermore, cathepsin D secretion by breast cancer cells has been shown to lead to matrix degradation via the formation of large acidic vesicles, which may facilitate matrix component digestion (Montcourrier et al. 1999). pS2 overexpression could be associated with an alteration in ECM deposition and increased cell mobility (Williams et al. 1996).

Genotoxic metabolites, such as catechol estrogens and quinone derivatives, arising from E2 catabolism are suspected to promote carcinogenesis via production of genotoxic metabolites. Indeed, CYP1B1 (cytochrome P450) hydroxylates E2 to the promutagenic metabolite 4-OH catechol estrogen (4-OH-E2) (Hayes et al. 1996). E2-induced underexpression of TGFβ, described here, could increase CYP1B1 expression (Dohr et al. 1997) and thus enhance E2 catabolism. On the other hand, our data show that E2 induces underexpression of UDP glucuronosyltransferase, which is involved in the catechol-quinone detoxification system (Rafiojanis et al. 2000), thus contributing to increased production of semiquinone (E2-3,4-semiquinone) and quinone (E2-3,4-quinone) from 4-OH-E2 (Fig. 4). However, quinones can be recycled to catechols in the presence of NAD(P)H menadione oxidoreductase (Long & Jaiwaal 2000). Consequently, underexpression of this gene (Table 2) and decrease of its activity lead to increased levels of toxic E2-3,4-derived quinones. Quinones can also be synthesized from ubiquinol conjugates in the presence of NAD(P)H dehydrogenase ubiquinone (Hatefi et al. 1985).

Overexpression of this gene, in E2-treated MCF7, reinforces the accumulation of quinone metabolites promoting tumor progression (Yager & Liehr 1996). Moreover, E2 decreases both NAD(P)H:quinone oxidoreductase activity and thiol-specific antioxidant mRNA level (Table 3), an effect which could result in free radical damage (Yim et al. 1994, Ernster et al. 1995, Montano & Katzenellenbogen 1997). Thus, E2 appears to play a further role in breast cancer development by modulating the expression of five enzymes involved in the control of E2-dependent genotoxic metabolites and free radicals production, thereby increasing the probability that mutations occur during DNA synthesis. In conclusion, E2 appears to promote breast cancer progression by stimulating cell growth, anchorage junctions remodeling and ECM disassembly, together with genotoxic metabolite accumulation. The slight changes in gene expression observed by microarray analyses (Tables 2 and 3) appear sufficient to promote tumor progression and invasiveness.

TAM, an E2 antagonist, reverses the effects of E2 upon expression of genes involved in genotoxic metabolite accumulation and cytoskeletal ECM remodeling in addition to the overexpression of profilin, which is known to promote actin filament assembly at the barbed end (Kang et al. 1999) and of HSD17 β4, which is known to inactivate E2 by conversion to estrone (Luu-The 2001) (Tables 2 and 3). This could explain, at least in part, how TAM can abrogate the E2-induced phenotype. The fact that cell-cycle progression was not inhibited can be explained by recent findings that TAM can function as a molecular agonist inducing cell-cycle-associated gene in breast cancer (Hodges et al. 2003). Interestingly, TAM treatment also affects the expression of several genes not regulated by E2, such as profilin, DDB-2, XR-CC1, Hsp90, REA, immunophilin, ERK1, HSD17 β4 and TGF-α. Some of these genes could be involved in TAM resistance. Overexpression of REA (Table 3), which competes with the coactivator SRC-1 in binding to ER and inhibits ER transcriptional activity (Delage-Mourroux et al. 2000), could explain the reversal of E2-dependent transcriptional activities observed with TAM treatment. Although antiestrogenic treatment increases the life expectancy of breast cancer patients, resistance and relapse phenomena can frequently occur (Johnston 1997). Hypotheses of TAM resistance include the following: 1. ER structural and functional modifications (Chander et al. 1993); 2. post-receptor interaction modifications; 3. paracrine secretion modifications (Katzzenellenbogen 1991); 4. pharmacologic changes (Osborne et al. 1992). Our results support the first hypothesis, since we observed, in MCF7 cells treated with TAM, decreases in expression of Hsp90 and immunophilin, both of which are required for the formation of functional ER heterocooperates capable of interacting with hormones (Schiene-Fischer & Yu 2001).

We showed here that TAM decreases the expression of TGFα (Table 3). Throughout the duration of TAM treatment, the inhibition of growth factor expression would promote selection of cells endowed with constitutively high expression of one or more growth factors or growth factor receptor (Wakeling 1990). Indeed, TGFα is constitutively expressed in many estrogen-independent cells (Bates et al. 1988). Evidence now shows that constitutive activity of growth factor can bypass the cell’s dependence on estrogen and provide a mechanism for hormone-independent growth (Johnston et al. 1992), thus making TAM treatment inefficient. This could explain, at least in part, the arising of TAM
resistance. Moreover, TAM increases ERK1 gene expression (Table 3), which could in turn enhance cell proliferation and inhibit TGFβ-induced antiproliferative responses via SMAD 2/3 phosphorylation (Kretzschmar 2000). This may indicate a paradoxical effect of TAM, explaining at least in part, the phenomenon of TAM resistance (Rabenoelina et al. 2002). Finally, the downregulating effects of TAM treatment upon two genes involved in the base excision DNA repair system, DDB-2 (Itoh et al. 1999, Nichols et al. 2000) and XRCCL1 (Miller et al. 2001), could lead to increased mutation rate in tumor cells.

In conclusion, our experimental approach allowed us to identify 32 early E2-induced genes, including 19 new genes. It also allowed us to detect the TAM-induced expression reversal of 11 E2-induced genes. In addition, we were able to report TAM-specific regulation of nine genes, including eight new ones. Integrative data analyses provided a global understanding of the cellular mechanisms alterations associated with E2 and TAM treatments. This knowledge could be important to predict cell behavior associated with antiestrogen treatment and subsequently determine new therapeutic targets. In particular, an understanding of mechanisms leading to TAM resistance/relapse may facilitate the development of novel therapeutic strategies addressed to estrogen-regulated tumorigenesis.

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