In search of novel mechanisms leading to the development of antiestrogen-resistance in human breast tumors, we analyzed differences in the gene and protein expression pattern of the human breast carcinoma cell line T47D and its derivative T47D-r, which is resistant toward the pure antiestrogen ZM 182780 (Faslodex™, fulvestrant). Affymetrix DNA chip hybridizations on the commercially available HuGeneFL and Hu95A arrays were carried out in parallel to the proteomics analysis where the total cellular protein content of T47D or T47D-r was separated on two-dimensional gels. Thirty-eight proteins were found to be reproducibly up- or down-regulated more than 2-fold in T47D-r versus T47D in the proteomics analysis. Comparison with differential mRNA analysis revealed that 19 of these were up- or down-regulated in parallel with the corresponding mRNA molecules, among which are the protease cathepsin D, the GTPases Rab11a and MxA, and the secreted protein hAG-2. For 11 proteins, the corresponding mRNA was not found to be differentially expressed, and for eight proteins an inverse regulation was found at the mRNA level. In summary, mRNA expression data, when combined with proteomic information, provide a more detailed picture of how breast cancer cells are altered in their antiestrogen-resistant compared with the antiestrogen-sensitive state.
The screening of global mRNA changes by hybridization of RNA to either cDNA or oligonucleotide arrays in a high-throughput manner has only become possible in recent years (9) and has been applied to the analysis of gene expression profiles of different breast cancer cell lines, to the study of the response of breast cancer cells to E$_2$, to the molecular profiling of breast cancer biopsies in relation to the ER$_x$ status, and to the successful subclassification of human breast tumors (10–15).

Proteomics is a complementary tool to mRNA analysis for assessing global changes in cellular protein expression and it can provide additional insight into post-translational modifications of proteins. Recent progress in the field of proteomics has resulted in publications on proteomes for normal and tumor-derived breast cancer cell lines and breast cancer biopsies, and results have been reviewed and discussed (16–21). Two-dimensional (2D) gel electrophoresis coupled with mass spectrometry (MS) identification of proteins has for example been carried out for establishing a catalogue of proteins that are expressed in human tumorigenic and nontumorigenic breast cell lines (22–27). Although cultivated cells tend to have a lower complexity of expressed proteins than human biopsies, comparison of only distantly related or rather unrelated cellular systems resulted in a large number of differentially expressed proteins, however with limited clinical relevance. The proteomics studies of normal human breast, premalignant lesions, and breast cancer biopsies generated data with stronger clinical impact, but even higher biological complexity (28–32). Moreover, in specialized approaches, luminal and myoepithelial breast cells from reduction mammoplasties were analyzed by proteomics after a double antibody magnetic sorting technique (33), and membrane proteins of breast cancer cell lines were studied after cell fractionation (34). This lowered the complexity of the systems and increased the number of low-abundance proteins that could be detected. Recently, a proteomics study of the human breast cancer cell line MCF-7 with or without treatment of the antiestrogen ZM 182780-resistant T47D-r has been reported (35).

Today, only a limited number of studies have been published in which both technologies, proteomics and mRNA expression profiling, have been compared using the same samples (36–38). There are reports that absolute protein and mRNA expression levels do not correlate well when total expression profiles are analyzed (39). With regard to a comparison of those genes and proteins that are differentially expressed, several studies have been published recently (40–47). The conclusion drawn by the authors on the correlation of differential RNA and protein expression varies between moderate and no correlation and seems to depend at least partially on the abundance of the mRNA or the protein under investigation.

The aim of the present study was to define and compare differential gene and protein expression in a model system of antiestrogen-sensitive and -resistant breast cancer and to evaluate concordance of mRNA and protein data. It is expected that a more detailed analysis of these changes will give new hints about the mechanisms that lead to the development of antiestrogen-resistant breast cancer and may eventually help to propose novel strategies and targets for an effective breast cancer therapy.

**EXPERIMENTAL PROCEDURES**

**Chemicals**—The steroid hormone E$_2$ and ZM 182780, synthesized in the Laboratories of Schering AG (Berlin, Germany), were solubilized in absolute ethanol at a stock concentration of 10$^{-3}$ and 10$^{-2}$ M, respectively, and stored at −20 °C. Dilutions were performed in absolute ethanol.

**Cell Culture**—The human breast cancer cell line T47D was obtained from the American Type Cell Culture Collection (Manassas, VA). The cells were routinely grown in phenol red-free RPMI 1640 medium (Biochrom, Berlin, Germany) supplemented with 10% fetal calf serum (LifeTechnologies, Karlsruhe, Germany), 2 mM glutamine (LifeTechnologies), 200 mM/ml insulin (Sigma, Deisenhofen, Germany), and 10$^{-10}$ M E$_2$. The parental cells were cultivated in parallel to the antiestrogen-resistant cells. Prior to experiments, the cells were exposed to standard medium: phenol red-free RPMI 1640 medium, supplemented with 10% charcoal-treated fetal calf serum, 2 mM glutamine, and 10$^{-10}$ M E$_2$. The antiestrogen-resistant cells were grown continuously in standard medium with 10$^{-6}$ M ZM 182780. Cells were grown at 37 °C in a humidified atmosphere with 5% CO$_2$. For generation of protein extracts, 8 × 10$^6$ T47D cells were seeded in standard medium and T47D-r in standard medium plus 10$^{-6}$ M ZM 182780 (day 0). On day 3, medium was replaced for 24 h with standard medium in the case of T47D and standard medium plus 10$^{-7}$ M ZM 182780 in the case of T47D-r. Cell cultures that had reached 70% confluence were harvested on day 4.

For isolation of RNA, T47D were cultivated in standard medium and T47D-r in standard medium plus 10$^{-7}$ M ZM 182780 for 24 h. Poly(A)$^+$ RNA from two independent cell culture experiments was hybridized to the HuGeneFL array, and total RNA from a further independent cell culture experiment was hybridized to the Hu95A array.

**Proliferation Assay**—The proliferation rates of ZM 182780-sensitive T47D and ZM 182780-resistant T47D-r were measured with the 4,5-dimethylthiazol-2-yl-2,5-diphenyltetrazolium (MTT) assay. Cells grown to 70–80% confluence were harvested and seeded in 200 μl in 96-well microtiter plates at 2500 cells/well in phenol red-free RPMI 1640, 2 mM glutamine, 10% charcoal-treated fetal calf serum (day 0). E$_2$ (10$^{-10}$ M), or the vehicle control ethanol at 0.001%, or E$_2$ (10$^{-10}$ M) plus ZM 182780 at concentrations of 10$^{-10}$ to 10$^{-6}$ M were added. Mitochondrial activity was measured on day 7 after plating by adding 20 μl MTT reagent (Sigma, 5 mg/ml in PBS) for 3–5 h at 37 °C to the medium. Afterward, the supernatant was removed and the formazan crystals were dissolved in dimethylsulfoxide. Absorbance was measured at 550 nm. Experiments were performed at least twice and one representative experiment with mean from 6 wells is shown.

**Generation of Protein Extracts for Proteomics Analysis**—For sample preparation, 1 × 10$^7$–1 × 10$^8$ cells were washed three times with PBS without Ca$^{2+}$, Mg$^{2+}$ to remove serum proteins and detached by treatment with PBS (–Ca$^{2+}$, –Mg$^{2+}$) containing 2 mM EDTA. The pelleted cells were resuspended in PBS containing a protease inhibitor mixture (Complete$^\text{TM}$ Mini EDTA-free, pepstatin, and phosphoramidon) (all from Roche, Mannheim, Germany) and sodium-orthovanadate (Sigma) and stored in aliquots as dry pellets at −80 °C before use. Cell pellets were resuspended in 20 mM Tris/HCl, pH 8.0, 2 mM MgCl$_2$ containing protease inhibitor mixture and sodium-orthovanadate and lysed at 4 °C with ultrasound (Sonorex RK31; Bandelin, Berlin, Germany) in the presence of glass beads of 1.7–2.0 mm diameter (Roth, Karlsruhe, Germany). The DNA in the lysate was digested with benzoylase (Merck, Darmstadt, Germany) at room temperature for 30 min.

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Afterward, lysis buffer was added containing 7 M urea (Amersham Biosciences, Uppsala, Sweden), 2 M thiourea (Merck), 1% (w/v) dithiothreitol (Sigma), 2% (w/v) 3-[(3-cholamidopropyl)dimethylammonio]-1-propanesulfonate (Electrophoresis Reagent; Sigma) and 0.5–1% immobilized pH gradient (IPG) buffer (Amersham Biosciences), corresponding to the pH gradient. The lysate was vortexed for 1 min and centrifuged. The protein concentration was determined in the supernatant with the modified Bio-Rad protein assay kit (Bio-Rad, Hemel Hempstead, UK) and was 3–5 µg/µl.

2D-SDS-PAGE—Samples were loaded onto IPG strips (Immobiline DryStrips; Amersham Biosciences) using 100 µg total protein for analytical gels, which were silver-stained, and 1500 µg total protein for preparative gels, which were stained with colloidal Coomassie brilliant blue G250 (Serva, Heidelberg, Germany). For the analytical gels, it was taken care that the amount of protein loaded onto the IPG strips was within the linear range of the silver stain detection. Spots that were saturated have not been included in the analysis. Isoelectric focusing was performed on an IPGphor unit (Amersham Biosciences) for IPG gradients pH 3–10, pH 5–8 (Bio-Rad), and pH 4–7 and on a Multiphor unit (Amersham Biosciences) for IPG gradients pH 6–11.

For the second-dimension SDS-PAGE using the Hoefer ISO-DALT system (Amersham Biosciences), focused Immobiline DryStrips containing traces of bromophenol blue (Amersham Biosciences) were run in SDS-PAGE running buffer (25 mM Tris, 192 mM glycine, 0.1% SDS) at 20 °C until the dye-front reached the end of the gel. All samples were run at least in duplicates from three independent cell culture experiments leading to 12 gels for each pH gradient, which were run in SDS-PAGE running buffer (30 mM Tris base, 40 mM 6-amino hexane acid, 20% methanol), anode I buffer (30 mM Tris base, 50 mM NaCl, 5 mM EDTA, 30 mM sodium-pyrophosphate, 50 mM NaF, 1 mM phenylmethylsulfonyl fluoride (Sigma), and the protease inhibitor mixture Complete™ Mini (Roche), phosphatase inhibitor cocktail I and II (Sigma), and 1% (w/v) Triton X-100 (Sigma). The plates were left on ice for 10 min, and the lysates were transferred to Eppendorf tubes. The lysate was vortexed 2 × 30 s and cell lysis was pelleted at 14,000 rpm for 20 min at 4 °C. The supernatant was frozen at –80 °C. Protein concentration of the cell lysate was determined with the DC Protein Assay Kit II (Bio-Rad). Equal amounts of protein were separated on 10% bis-Tris gels (NuPAGE®; Invitrogen, Karlsruhe, Germany) and blotted to Optitran BA-S 85 reinforced nitrocellulose membrane (Schleicher & Schuell, Dassel, Germany) by semi-dry blotting in cathode buffer (25 mM Tris base, 40 mM 6-amino hexane acid, 20% methanol), anode I buffer (30 mM Tris base, 20% methanol), and anode II buffer (300 mM Tris base, 20% methanol) at 0.8 mA/cm² for 60 min. The membrane was incubated in blocking buffer, i.e. Roti-Block (Carl Roth GmbH, Karlsruhe, Germany) for 90 min at room temperature. Incubation with the primary antibodies was carried out in blocking buffer at 4 °C overnight. The mouse monoclonal α-cathepsin D antibody Ab-1 (clone BC011) was purchased from Oncogene™ Research Products (Boston, MA), and the α-β-actin was a rabbit polyclonal serum purchased from Sigma. As secondary antibodies α-mouse and α-rabbit horseradish peroxidase-coupled antibodies (Amersham Biosciences) were used. The α-cathepsin D antibody was diluted 1:500, and the α-β-actin serum 1:1000 in blocking buffer. Afterward, membranes were washed 3 × 5 min in PBS-T (PBS with 0.1% Tween-20) and incubated with the respective secondary horseradish peroxidase-coupled antibody (diluted 1:2000) for 1 h at room temperature. Membranes were again washed 3 × 5 min in PBS-T. Detection was carried out with the ECL Plus system (Amersham Biosciences) according to manufacturer’s instructions. Blots were quantitated with the ImageJ software (National Institutes of Health, USA).

Affymetrix DNA Chip Hybridization—One to 2 µg of poly(A)⁺ RNA or 10 µg of total RNA were converted into double-stranded cDNA using a modified oligo-dT primer including a 5’ T7 RNA polymerase promoter sequence and the Superscript Choice system for cDNA synthesis (Life Technologies, Rockville, MD). In vitro transcription was performed with T7 RNA polymerase (T7 Megascript kit; Ambion, Austin, TX) and 0.5–1 µg of double-stranded cDNA template in the presence of a mixture of ATP, CTP, GTP, biotin-11-CTP, and biotin-16-UTP (ENZO Diagnostics, Farmingdale, NY). Twenty micrograms of cRNA were fragmented randomly by incubating in 40 mM Tris acetate, pH 8.1, 100 mM K⁺ acetate, and 30 mM Mg²⁺ acetate at 94 °C for 35 min. The Affymetrix HuGeneFL array and the Hu95A array were hybridized, washed, and stained according to standard protocols. The
the experiment. For each such pair-wise normalization, all genes that are the remaining arrays is normalized separately against the reference intensity of all arrays is chosen as the reference experiment. Each of the arrays is sorted by their intensity (median of all individual oligonucleotide signal intensities). The experiment with the median first step, all arrays are sorted by their intensity (median of all individual oligonucleotide signal intensities). The experiment with the median of all arrays is chosen as the reference experiment. Each of the remaining arrays is normalized separately against the reference experiment. For each such pair-wise normalization, all genes that are present on both arrays are identified. A gene is considered present if the p value for a one-sided matched-pairs signed-rank Wilcoxon test applied to all unmasked perfect match (PM) versus mismatch (MM) oligonucleotide signal intensities is below 0.04. To identify a set of oligo probe pairs belonging to genes with unchanged expression, all probe pairs for genes present on both arrays are sorted by their intensity differences of perfect match minus mismatch oligonucleotide (PM-MM) and are assigned the corresponding ranks. That is, each probe pair has a rank on the reference array and a usually different rank on the array to be normalized. Now, an iterative process is used to remove probe pairs with high rank jumps between the two experiments until a subset of probe pairs with identical ranks on both arrays is obtained (maximum order preserving set). The assumption is that probe pairs with high rank jumps belong to genes that do change their expression, whereas probe pairs with small rank jumps belong to genes with constant expression. For all probe pairs from the order preserving set, the intensity difference PM-MM on the reference array is plotted against PM-MM on the array to be normalized and the best-fitting forth order polynomial with no zero-order term is identified. In a last step, all PM-MM differences from the array to be normalized are transformed by the thus identified nonlinear function. In case of the HuGeneFL array, where two independent hybridization experiments have been performed, the average expression change between the replica experiments had to be at least 2-fold to be considered as regulated.

RESULTS

Characterization of T47D and T47D-r—T47D-r were generated from the antiestrogen-sensitive human breast cancer cell line T47D after continuous cultivation in presence of \(10^{-10}\) M E\(_2\) and \(10^{-6}\) M antiestrogen ZM 182780 for 31 weeks (8). Phase contrast photographs of T47D and T47D-r cells cultivated for 3 days in standard growth medium are shown (Fig. 1). Parental T47D and the ZM 182780-resistant derivative T47D-r have a similar morphology that is typical of tumor cells with an apolarized polygonal morphology.

The proliferative response of T47D and T47D-r to E\(_2\) and increasing concentrations of the antiestrogen ZM 182780 was measured with the MTT assay (Fig. 2). Cells were seeded in 96-well plates and immediately treated with ethanol, or \(10^{-10}\) M E\(_2\), or \(10^{-10}\) M E\(_2\) plus ZM 182789 at concentrations from \(10^{-10}\) M to \(10^{-6}\) M. Whereas T47D are growth-inhibited by increasing concentrations of ZM 182780, the antiestrogen-resistant T47D-r grow equally well in medium with vehicle ethanol or increasing concentrations of ZM 182780.

Differentially Expressed Genes—RNA from T47D and T47D-r was hybridized to the Affymetrix arrays HuGeneFL and Hu95A. The results from Affymetrix DNA chip hybridization are displayed in Table I. Out of a total of 6574 probe sets that interrogate genes and ESTs and that are represented on the HuGeneFL array, 104 genes were found consistently up-regulated and 63 down-regulated more than 2-fold in two independent cell culture experiments in T47D-r versus T47D cells. Out of a total of 12,387 probe sets on the Hu95A array, 231 genes were up-regulated and 316 genes down-regulated more than 2-fold when RNA from a single cell culture experiment was hybridized. To determine the number of genes that are interrogated more than once on the two different Affymetrix arrays we counted the number of probe sets that are represented as singletons or are members of a cluster resulting in a nonredundant gene data set. First, from the total of 18,961 probe sets present on both chips, 91 probe sets had to be removed because they contained low complexity gene information. The remaining 18,870 probe sets fall into two groups with 3,777 singletons and 7,142 clusters containing...
Fig. 2. Response of T47D and T47D-r to E₂ and the antiestrogen ZM 182780. Cell proliferation was measured with the MTT assay. Cells were seeded in 96-well plates and immediately treated with ethanol or 10⁻¹⁰ M E₂, or 10⁻⁸ M E₂ plus ZM 182780 at concentrations from 10⁻¹⁰ m to 10⁻¹⁶ m. Mitochondrial activity was measured on day 7. Proliferation activity for E₂-treated T47D and T47D-r, respectively, was set to 100. Whereas T47D are growth-inhibited by increasing concentrations of ZM 182780, the antiestrogen-resistant T47D-r grow equally well in medium with vehicle ethanol or increasing concentrations of ZM 182780.

| Array              | Probe sets | Up-regulated | Down-regulated | Total |
|--------------------|------------|--------------|----------------|-------|
| HuGeneFL           | 6574       | 104          | 63             | 167   |
| Hu95A              | 12387      | 231          | 316            | 547   |
| HuGeneFL + Hu95A   | 18961      | 335          | 379            | 714   |
| Nonredundant genes | 10919      | 266          | 300            | 566   |

Table I
Overview of genes differentially expressed in T47D-r versus T47D analyzed on Affymetrix chips HuGeneFL and Hu95A

On the HuGeneFL array RNA from two independent cell culture experiments and on the Hu95A array RNA from one further independent cell culture experiment was hybridized. Genes that are regulated more than 2-fold on the arrays in T47D-r and T47D are listed.

15,093 sequences that result in 10,919 nonredundant genes (Table I). The same type of analysis was performed for the up- and down-regulated probe sets on the two array types. Taken together, 266 nonredundant genes are up-regulated and 300 are down-regulated in T47D-r compared with T47D (Table I). An in-depth analysis of the Affymetrix DNA chip hybridization results will be described elsewhere.²

2D-PAGE and Image Analysis—In order to obtain well-distributed spot patterns on overview gels (pH 3–10), the lysing conditions for the human breast cancer cell line T47D and its derivative T47D-r were optimized. Reproducible protein expression patterns were observed with the majority of proteins visible between pH 5–7. To increase the resolution of the gels within this pH range and to obtain a higher number of well-separated spots, the samples from three independent cell culture experiments were analyzed on narrow pH gradients (pH 4–7 and pH 6–11) (Fig. 3). For image comparison, the gels were scanned and automatic spot detection was performed with the Phoretix-2D advanced software. Depending on the spot detection parameters, manual editing of the images was found to be necessary mainly to split spots, to erase false spots, and to include missed spots. These efforts resulted in ~2500 spots on the gels. After matching the gels derived from the T47D cell line with the gels from its derivative T47D-r, many spots appeared to be variant by at least a factor of two in spot volume (Figs. 3 and 4). In the pH range 4–7 53 proteins and in the pH range 6–11 another 31 proteins were differentially expressed. Of these 84 proteins, 46 were consistently up-regulated and 38 down-regulated in the three independent cell culture experiments of T47D-r versus T47D.

Identification of Regulated Proteins—Proteomics analysis identified 84 protein spots that were consistently elevated 2-fold or more in three independent cell culture experiments of T47D or T47D-r (Fig. 5). Of these, 73 differential protein spots were excised from preparative Coomassie-stained 2D gels. As the amount of protein loaded on preparative Coomassie-stained gels is on the upper limit of the capacity of the IPG strips, we established a further pH gradient (pH 5–8) to get a good resolution in the overlap region of the pH gradients pH 4–7 and pH 6–11. The excised spots were subjected to trypsin digestion, and resulting peptide fragments were analyzed by MS. Assignments have been obtained for 66 out of 73 spots. Among the 66 identified protein spots, 11 proteins were found in more than one spot, e.g. cathepsin D was identified in six independent spots (Fig. 4, Table II) and MxA in two independent spots (Table II). As a consequence, 47 distinct proteins were identified. To confirm differential protein expression of cathepsin D, a Western blot analysis was performed (Fig. 6), and the blot was quantitated after normalization to β-actin. Cathepsin D is down-regulated 2.4-fold in T47D-r treated with E₂ and ZM 182780 compared with T47D treated with E₂.

Differentially Expressed Proteins—The differentially regulated proteins belong to several functional categories (Tables II and III). Thirteen proteins are enzymes and for seven the function is unknown. Six signal transduction proteins and six proteins with a role in protein degradation were identified. Four proteins are of mitochondrial origin, two each have a function in the cytoskeleton, are secreted proteins, or are involved in transformation. Of the 47 proteins, 44 were either up- or down-regulated, whereas three proteins (stathmin/oncoprotein 18, 14-3-3 γ, and Nudix hydrolase NUDT5), which each were identified from two or three independent spots on the gel, were regulated both up in one spot and down in

² A. Sommer, personal communication.
another, in T47D-r compared with T47D (Fig. 5). Concerning the 44 proteins regulated in only one direction, 20 proteins were up-regulated and 24 proteins were down-regulated in the comparison of T47D-r versus the T47D.

**Comparison of Proteomics and Affymetrix Data**—Only the 44 distinct proteins that were regulated up or down were compared with mRNA expression data (Fig. 5). To achieve this, the sequence of the protein that was identified by MS or the corresponding DNA sequence was used to perform a tBLASTn or BLASTn search, respectively, of the databases that contain the genes present on the Affymetrix arrays. This analysis showed that of the 44 proteins six were not present on the two arrays. Comparison of differential mRNA with protein expression data revealed that 19 out of 38 proteins for which the corresponding genes were present on the two arrays were regulated in the same direction on the mRNA and protein level, i.e. 50% of the proteins show a concordant regulation on the mRNA and protein level.

However, the strength of differential expression can vary. For 11 proteins the corresponding mRNAs are not well defined, i.e. either only one differential expression value is available and the fold-change is below 2 or the fold change is weak and not consistent between the experiments. Eight proteins were inversely regulated on the protein compared with the mRNA level. The differential expression of these mRNAs was often below 1.8-fold (Table II), whereas in standard Affymetrix experiments only genes regulated by a factor of two and higher are considered for subsequent studies.

**DISCUSSION**

**Comparison of Differential Proteomics and RNA Expression Analysis**—So far only a limited number of studies compared...
differential gene and protein expression (38, 40–47). In these studies, cancer and normal tissue, different cell lines, and treated and untreated cellular model systems (e.g. yeast) were compared. The correlation of differential RNA and protein expression varied between moderate and no correlation according to these reports. Poor correlation was seen in a study of human liver (39), whereas a moderate correlation was reported in yeast (40). One group reported that for well-resolved and abundant known proteins a highly significant correlation between protein levels as judged by 2D-PAGE and gene expression profiling analysis in noninvasive and invasive human transitional cell carcinomas was found (38). Several mRNAs and proteins showed a striking correspondence, although in some cases also discrepancies were seen that could potentially be attributed to mRNA stability, splicing, translational regulation, post-translational processing, control of protein turn-over, protein degradation, or a combination of these.

To extend previous studies, it was our aim to analyze differential gene expression on both the mRNA level by Affymetrix DNA chip hybridization and on the protein level by proteomics. In addition, we expected to achieve a better insight into the changes that accompany the development of antiestrogen-resistance in our system.

Affymetrix DNA chip hybridizations were carried out on the commercially available HuGeneFL and Hu95A array. In sum, a total of 10,919 nonredundant genes and ESTs were analyzed on the two arrays. A total of 266 genes (2.4%) were found to be up-regulated and 300 (2.7%) down-regulated more than 2-fold in T47D-r compared with T47D (Table I).

Compared with the proteomics analysis, where only 84 differentially regulated spots were identified on 2D gels, far more genes were found to be differentially expressed. This can be explained by mRNA measurements having on the one hand a higher degree of sensitivity and the capability to detect also weakly expressed mRNAs and on the other hand oligonucleotides arrayed in parallel allowing for a higher degree of automation and throughput.

When the total cellular protein content from three independent cell culture experiments of T47D and T47D-r was analyzed, 84 spots were found to be reproducibly variant more than 2-fold. Sixty-six proteins were identified corresponding to 47 distinct proteins because 11 proteins were present in more than one spot on the gel. The fact that one protein is localized in two different spots on the gel can be explained by post-translational modifications, e.g. phosphorylation, glycosylation, or limited proteolytic cleavage (48, 49).

Three proteins (stathmin/oncoprotein 18, 14-3-3 γ, and Nudix hydrolase NUDT5) were identified from two or three independent spots on the gel and were regulated both up in one spot and down in another in T47D-r compared with T47D (Fig. 4).
Comparison of Proteomic and Genomic Analysis of T47D and T47D-r

**TABLE II**

Proteomics data for all analyzed protein spots and comparison with corresponding mRNA expression

| Spot ID | pH gradient | Fold regulation T47D-r versus T47D | GenBank protein identifier | Identified protein | Proteomics M theor. | Proteomics M exp. | Proteomics pI | Proteomics total pep. in digest | Matched pept. / total pep. in digest | Error | seq. cov. | Gene symbol | GenBank nucleotide identifier | DNA chip hybridization |
|---------|-------------|------------------------------------|-----------------------------|--------------------|---------------------|-------------------|----------------|-------------------------------|-------------------------------------|-------|-----------|-------------|-----------------------------|-------------------------|
| 3       | pH 4–7      | 6.3                                | gi 12652557                 | Similar to actin-related protein 2/3 complex, subunit 5 | 17                 | 15                | 6.2            | 6.3                          | 4/5                  | 9     | 33        | NM_030978   | np                          | np | np |
| 6       | pH 4-7      | 1.5                                | gi 13569956                 | Unknown, clone 24952 | 19                 | 14                | 5.2            | 5.4                          | 7/10                  | 14    | 36        | AF131758    | np | np | np |
| 33      | pH 4-7      | –2                                 | gi 6018458                  | 6-phosphogluconolactase | 28                 | 25                | 5.7            | 5.9                          | 11/12                 | 30    | 49        | PGLS        | AJ243972     | np | np | np |
| 56      | pH 6–11 only in T47D-r | 10.75624 | gi 131528 | Poly(ADP-ribose) polymerase | 57                 | 58                | 9.3            | 8.7                          | 4/4                   | 22    | 8         | PTB2        | X66975      | np | np | np |
| 57      | pH 6–11     | 1.8                                | gi 1649555                  | Vacular sorting protein | 57                 | 58                | 9.3            | 8.7                          | 6/6                   | 11    | 12        | np | np | np |
| 67      | pH 5–8      | –3.3                                | gi 4680711                  | CIG-36 protein       | 19                 | 27                | 6.4            | 7.2                          | 9/11                  | 20    | 33        | NM_015963   | np | np | np |
| 73      | pH 5–8      | 3.9                                | gi 5450464                  | SYT interacting protein | 70                 | 80                | 9.7            | 8.7                          | 16/24                 | 16    | 26        | RBM14       | AF080561    | np | np | np |
| 53      | pH 5–8      | 2.2                                | gi 5450640                  | Keratin, type II cytoskeletal 4 | 58                 | 60                | 6.3            | 7.2                          | 9/14                  | 17    | 19        | KRT4        | X07695      | np | np | np |
| 56      | pH 6–11     | 2.2                                | gi 5450603                  | Protein phosphatase 1 alpha isoform | 38                 | 39                | 5.9            | 6.2                          | 16/24                 | 16    | 49        | PPPI1A      | S57501      | np | np | np |
| 47      | pH 6–11     | 2.9                                | gi 5450599                  | 4-nitrophenylphosphatase domain and non-neuronal SNAP2-1 like 1 | 33                 | 34                | 9.5            | 8.4                          | 12/14                 | 23    | 26        | NIPS1AP1    | AJ001258    | np | np | np |
| 48      | pH 6–11     | 2.7                                | gi 5450779                  | Ubiquitin-conjugating enzyme E2M | 21                 | 17                | 7.7            | 7.2                          | 7/8                   | 14    | 35        | UBE2M       | AF075999    | np | np | np |
| 49      | pH 6–11     | 2.4                               | gi 5450551                  | Peroxiredoxin 1, proliferation-associated gene A (natural killer-enhancing factor A) | 22                 | 19                | 8.7            | 7.7                          | 19/22                 | 20    | 64        | PRDX1       | X67951      | np | np |

**Inversely regulated on protein and mRNA level**

| 4       | pH 4–7      | –2.9                                | gi 238470                  | Acid phosphatase isozyme BF or AF | 18                 | 14                | 7.1            | 6.4                          | 12/14                 | 14    | 81        | ACPI        | U25849      | 1.4 | 1.7 | –1.1 |
| 27      | pH 4–7      | –2                                 | gi 4820738                 | Tumor protein D52 | 20                 | 22                | 4.9            | 5.0                          | 6/6                   | 14    | 38        | TP53        | U18914      | undef | 1.2 | 1.3 |
| 34      | pH 4–7      | –6.2                                | gi 1352435                 | Eukaryotic translation initiation factor 4E | 25                 | 25                | 5.8            | 6.0                          | 6/7                   | 31    | 22        | EIF4E       | M15353      | 1.1 | –1.1 | 1.5 |
| 45      | pH 6–11     | –2.3                                | gi 4507777                  | Ubiquitin-conjugating enzyme E2D 3 | 17                 | 12                | 8.0            | 7.3                          | 8/11                  | 9     | 34        | UBE2D3      | U39318      | 2.2 | 1.2 | 1.5 | 1.3 |
| 60      | pH 6–11 only in T47D-r | 10.75624 | gi 4504327 | Hydroxyacyl-Coenzyme A dehydrogenase | 52                 | 57                | 9.6            | 8.7                          | 9/10                  | 25    | 21        | HADHB       | D16481      | –1.2 | –1.2 | –1.4 |
| 61      | pH 6–11     | 5.1                                | gi 4758504                  | Hydroxyacyl-Coenzyme A dehydrogenase, type II | 52                 | 57                | 9.6            | 8.7                          | 15/18                 | 18    | 35        | HADHB2      | U73514      | <–1.6 | –1.5 | np |
| 62      | pH 6–11     | 6.2                                | gi 4758504                  | Hydroxyacyl-Coenzyme A dehydrogenase, type II | 52                 | 57                | 9.6            | 8.8                          | 14/15                 | 26    | 35        | HADHB2      | U73514      | np | np | np |
| 63      | pH 6–11     | 1.6                                | gi 4758504                  | Hydroxyacyl-Coenzyme A dehydrogenase, type II | 27                 | 22                | 8.0            | 7.1                          | 15/17                 | 20    | 75        | HADHB2      | U73514      | <–1.6 | –1.5 | np |
| 64      | pH 6–11     | 1.6                                | gi 4758504                  | Hydroxyacyl-Coenzyme A dehydrogenase, type II | 27                 | 21                | 8.0            | 7.1                          | 5/6                   | 20    | 17        | HADHB2      | U73514      | np | np | np |
| 65      | pH 6–11     | –2.5                                | gi 5902074                  | Stromal cell-derived factor 2 | 23                 | 21                | 7.2            | 7.1                          | 6/7                   | 31    | 35        | SDF2        | D50645      | 1.4 | –1.1 | –1 |
| 66      | pH 5–8      | –2.5                                | gi 4506195                  | Proteasome subunit, beta type, 2 | 23                 | 28                | 6.5            | 7.4                          | 13/18                 | 11    | 50        | PSMB2       | D25699      | 1.7 | 1.2 | 1.1 |

**Regulated in parallel on mRNA and protein level**

| 1       | pH 4–7      | –3.6                                | gi 599882                   | Cellular retinoic acid-binding protein II | 16                 | 11                | 5.4            | 5.6                          | 16/20                 | 14    | 82        | CRBP2       | M97815      | –2.5 | –2.8 | –1.6 |
| 7       | pH 4–7      | 1.5                                | gi 2392338                  | Glyoxalase I | 21                 | 20                | 5.1            | 5.0                          | 10/14                 | 14    | 51        | GLO1        | D13315      | 1.4 | 1.6 | 1.2 |
## Table II—continued

| Spot ID | pH gradient | Fold regulation T47D-r versus T47D | GenBank protein identifier | Identified protein | M<sub>th</sub>, M<sub>exp</sub> | pl theor, pl exp | Matched pepT/total pepT in digest | Error<sup>2</sup> | Gene symbol | GenBank nucleotide identifier | DNA chip hybridization |
|---------|-------------|------------------------------------|---------------------------|-------------------|----------------|------------------------|----------------|-------------|----------------------------|--------------------------|
| 9       | pH 4–7      | 1.6                                | gi 4502969                | Catechol-O-methyltransferase | 21/20 5.1 5.1 | 9/14 20 39 | COMT | Z26491 | 2.1 | M58525 | >2.3 | 1.9 | np |
| 10      | pH 4–7      | 1.5                                | gi 4557032                | Lactate dehydrogenase B | 20/36 6.1 5.9 | 8/12 21 14 42 | LDHB | X13794 | undef | >10 | 4.1 | 1.9 | np |
| 11      | pH 4–7      | 7.8                                | gi 4575804                | Serine protease inhibitor B | 27/27 5.3 4.7 | 11/14 28 17 | FAH  | M55150 | 1.1 | M20560 | >2.7 | 5.6 | np |
| 12      | pH 4–7      | 6                                  | gi 4550291                | Phosphomonoesterase 6 | 22/26 5.6 6.3 | 13/14 24 16 | YWHA  | AF010126 | 0.9 | M31303 | >2.7 | 5.6 | np |
| 13      | pH 4–7      | 2                                  | gi 4687601                | Bacterial protein        | 27/27 5.3 4.7 | 11/14 28 17 | FAH  | M55150 | 1.1 | M20560 | >2.7 | 5.6 | np |
| 14      | pH 4–7      | 2                                  | gi 4553545                | Antigen related protein 2 | 20/27 5.6 5.8 | 11/14 28 17 | FAH  | M55150 | 1.1 | M20560 | >2.7 | 5.6 | np |
| 15      | pH 4–7      | 2                                  | gi 4557587                | Fumarase coreactase      | 27/27 5.3 4.7 | 11/14 28 17 | FAH  | M55150 | 1.1 | M20560 | >2.7 | 5.6 | np |
| 16      | pH 4–7      | 2                                  | gi 4550291                | Phosphomonoesterase 6 | 22/26 5.6 6.3 | 13/14 24 16 | YWHA  | AF010126 | 0.9 | M31303 | >2.7 | 5.6 | np |
| 17      | pH 4–7      | 2                                  | gi 4553545                | Antigen related protein 2 | 20/27 5.6 5.8 | 11/14 28 17 | FAH  | M55150 | 1.1 | M20560 | >2.7 | 5.6 | np |
| 18      | pH 4–7      | 2                                  | gi 4557587                | Fumarase coreactase      | 27/27 5.3 4.7 | 11/14 28 17 | FAH  | M55150 | 1.1 | M20560 | >2.7 | 5.6 | np |

1. Autolytic tryptic peptides and contaminants were not taken into account.
2. Average error of matched peptide masses.
3. Two peptides confirmed by electrospray ionization MS/MS.
4. np, Not present on array.
5. One peptide confirmed by MALDI-post-source decay.
6. Spot saturated.
7. undef, Undefined, indicating that in the two compared chip hybridizations no signal was detected.
8. One peptide confirmed by electrospray ionization MS/MS.
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5). The regulation in different directions in separate spots could also be the result of post-translational modifications. For instance, for 14-3-3 γ and stathmin signal-induced phosphorylation has been described (50, 51). These three distinct proteins were excluded from the comparison with the Affymetrix results because their total amount was not investigated in this experimental setup. Another six proteins, which were not present on the two arrays hybridized, were also not included in the comparison with the Affymetrix results (Fig. 5).

The comparison of differential mRNA with protein expression data revealed that 19 out of 38 distinct proteins were up- or down-regulated in parallel at the mRNA and protein level, i.e. there is a concordance for the direction of changes on mRNA transcript and protein expression level for 50% of the distinct proteins for which a probe set was present on the two arrays.

In 11 cases, the differential expression level was not well-defined, therefore no comparison between protein and mRNA regulation could be made. For eight proteins, an inverse regulation on protein versus mRNA level was observed. This inconsistency between differential mRNA transcript and protein expression levels in a steady-state system, i.e. antiestrogen-sensitive and antiestrogen-resistant human breast cancer cells, as investigated here could be due to the presence of regulatory influences that are currently not understood (52). All explanations taken into account would require that mechanisms as detailed below have to work differentially between the parental cell and its antiestrogen-resistant derivative. In case the protein is up-regulated and mRNA is unchanged or down-regulated, one could postulate that either this particular protein is translated with higher efficiency, or it has a longer half-life, or that due to post-translational modification the protein appears de novo in an independent spot. When the protein is down-regulated while the mRNA is unchanged or up-regulated, the protein could have a shorter half-life, or it is translated with a lower efficiency. Further possibilities are that post-translational modification mechanism are turned off or that the protein is degraded into products no longer detectable because of various technical reasons (e.g. molecular weight or detection limit) leading to disappearance or decreasing intensity of this particular spot.

Because only those spots were considered to be differentially expressed in the proteomics analysis that were regulated by a factor of two or more, it is conceivable that a given protein is present in further spots for reasons described above but was not detected as being regulated or expressed below the detection limit. If the identity of all further possible isoforms of a protein has not been analyzed completely, this may influence the correlation between mRNA and protein and represents a limitation. In addition, when one looks in detail at the fold regulation on the mRNA level these factors are often below 1.8. In a standard gene expression analysis, only mRNAs with factors ≥2-fold are considered for subsequent analyses. A comparison of individual protein quantities from 2D gels with mRNA expression data has therefore to be performed with caution.

Biological Interpretation of the Data—In order to investigate the identified proteins further, we sorted them according to their functional category (Table III). Several of them are housekeeping enzymes, mitochondrial proteins, or involved in protein degradation. However, we also found proteins that play a role in signaling pathways or whose function is currently unknown. For several proteins that were identified in the proteomics analysis and by DNA chip hybridization, their role in regard to regulation by E2 and endocrine resistance will be depicted.

The protease cathepsin D was down-regulated on the protein and mRNA level in T47D-r compared with T47D. The protein was found at six different locations at the same apparent molecular mass (33 kDa) but different isoelectric points (between pH 5 and 6.5) on the 2D gels (Fig. 4). The different locations indicate that isoforms of cathepsin D exist. Cathepsin D is synthesized as pre-proenzyme that is post-transla-
tionally phosphorylated (53, 54). The 52-kDa pro-cathepsin D is cleaved to the enzymatically active 48-kDa heterodimer consisting of a 14- and 34-kDa subunit, and only the latter was identified in this proteomics study. The fold change on the protein level varied for the six spots between undetectable in T47D-r compared with T47D and down-regulated 2.9- to 2.1-fold (Table II). In the Western blot analysis, the cathepsin D form of 32 kDa was down-regulated 2.4-fold in T47D-r compared with T47D (Fig. 6).

On the mRNA level cathepsin D was down-regulated by a factor of more than 7.5- and 4.3-fold on the HuGeneFL array in T47D-r compared with T47D in the first and second cell culture experiment, respectively, and in the third experiment on the Hu95A array cathepsin D was down-regulated 7.3-fold.

Already in an early proteomics study on breast cancer cell lines a secreted 46-kDa protein was identified that was induced by E2 in several human ERα-positive cell lines and later determined by specific antibodies to be the protease cathepsin D (55). Thus, the cathepsin D gene is a well-known E2-regulated gene, and induction by E2 has been demonstrated for the ERα-positive breast cancer cell lines T47D, ZR-75-1, and MCF-7 (56, 57). Because T47D-r cells have lost expression of ERα, the E2-induction of cathepsin D is abrogated when compared with T47D cells. Differential expression of cathepsin D was found in both human breast cancer cell lines, breast biopsies, and reduction mammoplasties (33, 35).

The small GTPase Rab11a was down-regulated in T47D-r compared with T47D on the protein and mRNA level. On the protein level, Rab11a was found to be decreased 6.6-fold in T47D-r versus T47D. In the HuGeneFL and Hu95A array hybridization experiments, Rab11a was down-regulated by a factor of 1.8, 2.8, and 2.3 in mRNA from three independent cell culture experiments. Rab11a belongs to the large family of Ras-related small GTP-binding proteins that are involved in the regulation of vesicular transport along the endocytotic and exocytotic pathway (58, 59). It has been shown that Rab11a is regulated by E2 in the human endometrial cell line Ishikawa after transfection with either ERα or β, whereas treatment with ZM 182780 inhibited the E2-dependent induction of Rab11a (60). The fact that Rab11a is down-regulated on the protein and mRNA level in the breast carcinoma cell line T47D-r, which is continuously treated with ZM 182780 and has lost ERα expression, indicates that also in this cell line expression of Rab11a might at least be partially under the control of the ER. Interestingly, the Rab11a protein was recently found to be up-regulated in matched pairs of microdissected ductal carcinoma in situ compared with normal ductal breast tissue by an extensive 2D-PAGE-based proteomics analysis, and results were confirmed by immunohistochemical staining (32).

Another GTPase, the MxA protein, was identified from two responsive, well-differentiated breast cancer (66). As the T47D-r cells have lost protein and mRNA expression of the ERα (8), the identification of hAG-2 protein in the ER-positive T47D cells confirms the results shown previously on the mRNA level. Although the function of hAG-2 in the mammalian system is unknown, based on the expression data it was proposed that hAG-2 might be involved in the tumor cell biology of hormone-responsive, well-differentiated breast cancer (66).

In a proteomics study in which the plasma membranes of several ERα and epidermal growth factor receptor-positive human breast cancer cell lines were enriched, the hAG-2 homologous protein hAG-3 was identified (34), and in a focused follow-up study, hAG-2 and hAG-3 were analyzed by immunohistochemical methods (67). Here, it was demonstrated by quantitative reverse transcriptase PCR and immunohistochemistry on tissue microarrays that hAG-2 expression correlated positively with ER and negatively with epidermal growth factor receptor expression (67). Bioinformatic analysis indicated that the hAG-2 promoter contains four putative estrogen response elements. In a yeast 2-hybrid screen, hAG-2 was shown to interact with α-dystroglycan (DAG-1) and glycosylphosphatidylinositol-anchored protein C4.4a (67), and it was suggested that hAG-2 as secreted protein interacts with the extracellular matrix.

Many other proteins were found to be variant in our model system, which have not yet been described in association with the development of breast cancer and which are now further investigated to understand their role in endocrine-resistant breast cancer.

Conclusion—In the present study, we identified and compared differentially expressed genes and proteins in a model system of antiestrogen-sensitive and -resistant breast cancer and evaluated consistency between mRNA and protein data. In our steady-state model, concordance was 50%. By combining proteomics and Affymetrix technologies, we achieved a more detailed insight into the expression changes that accompany the development of antiestrogen-resistance in an in...
vitro system. RNA expression profiling on Affymetrix oligonucleotide arrays is a complementary method to proteomics as gel-based proteomics is often unable to detect the low-abundance proteins. Several known E2-regulated genes like cathepsin D or genes co-expressed with the ERα and likely also E2-regulated like hAG-2 are expressed in the parental T47D cell line. As manifestation of the antiestrogen-resistant phenotype of the T47D-r cells cathepsin D and hAG-2 are downregulated on the RNA and protein level. The discovery of gene and protein expression changes in the context of antiestrogen resistance may allow for a better understanding of the mechanisms that lead to endocrine-resistant breast carcinoma.

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