

A new gene expression signature related to breast cancer Estrogen Receptor status

E. Christodoulou, M. Ioannou, M. Kafousi, E. Sanidas, G. Papagiannakis,
V. Danilatou, G. Tsiliki, T. Margaritis, H. Kondylakis, D. Manakanatas, L. Koumakis,
A. Kanterakis, S. Vassilaros, M. Tsiknakis, A. Analyti,
G. Potamias, D. Tsiftsis, E. Stathopoulos, D. Kafetzopoulos

Abstract—The aim of this study is to identify a gene signature which is characteristic of ER status in breast cancer patients. To our knowledge, this is the first microarray study in Greece concerning breast cancer. We identified 97 genes that are characteristic for ER status and can well distinguish the ER+ from the ER- samples. We also managed to shrink our list to a 11-gene list giving the same prognostic outcome. We found a significant overlap of these genes with published prognostic signatures like the ones of West et.al. [4] and of Van't Veer et. al. [1]. This fact is very encouraging given the minimal overlap of such genes reported by others [28]. In order to give a meaningful information to the doctors we found associations with biological pathways. Interestingly, the vast majority of these genes are highly related to breast cancer.

I. INTRODUCTION

Expression profiling is a relatively recent technology that has gained the respect of the scientific community and it is now increasingly used. Its effectiveness and accuracy may overcome other caveats such as complexity and cost. In the present work we used expression profiling of breast tissue samples examining especially the levels of Estrogen Receptor (ER) and related genes.

It is well known that ER status is a strong marker that distinguishes different pathologic subtypes of breast cancer with prognostic implications [1], [4]. ER is a protein found inside the cells of the female reproductive tissue, some other types of tissue, and some cancer cells. The hormone estrogen will bind to the receptors inside the cells and may cause the cells to grow [2]. Many microarray studies exist on breast cancer but few of them conclude in a prognostic of ER status **gene signature** [26]. In this study, we tried to find a set of genes whose expression values are highly correlated with ER expression and associate them with possible biological meaning, an information important for the medical doctors. To our knowledge, this is the first microarray study in Greece aiming at the definition of an ER-related gene signature. This means that we are very

This work was partially funded by EPAN (AP14394/4-11-03), AKMON (AP6260EFA1250/17-5-2004) and PEP (KR8/16/6/2006) projects

The authors are with 1) FORTH, Institute of Molecular Biology and Biotechnology and Institute of Computer Science, P.O. Box 1385, 711 10, Heraklion, Greece, 2)The University of Crete, Medical School, P.O. 2208, Voutes, 71003 Heraklion, Crete, 3)Prolipsis Diagnostic Breast Center, 3 Sevastias Street, 115 28 Athens, Greece
echristo@imbb.forth.gr, mioannou@imbb.forth.gr

interested in examining whether the Greek patient population can be categorized into ER positive and ER negative based on known genes or whether there are some specific to the population genes that regulate the estrogen receptor status. This attempt is part of the *Prognochip* project. The *Prognochip* project aims at the identification and validation of *signature* gene expression profiles of breast tumors correlating with other epidemiological or clinical parameters. Towards these goals scientists from distant scientific disciplines join forces and efforts: Molecular Biology (Institute of Molecular Biology and Biotechnology, FORTH; <http://www.imbb.forth.gr>), Medicine (University Hospital, University of Crete Surgical Oncology; and "Prolipsis" a diagnostic center in Athens), Biostatistics and Computer Science (Institute of Computer Science, FORTH; <http://www.ics.forth.gr>) [8]

II. MATERIALS AND METHODS

A. Samples

26 fresh frozen breast tumors (17 ER+ and 9 ER-) were collected from patients treated at University Hospital of Heraklion after institutional review board approval. However, in this study we used 20 out of 26 samples because we selected only the strongly expressed ones (> 70%) ER+ aiming at a clearest signature. We assessed the size of our sample using the R package *samr* and found that for a mean difference of $\log_2 2$ (between the two groups) the False Discovery Rate (FDR) is 0.05. The False Negative Rate (FNR) is very low.

B. Array fabrication

The human library was obtained from the Qiagen and contains 34,580 70mer probes representing 24,650 genes and 37,123 gene transcripts. The lyophilized oligo set was resuspended in 3x SSC, 5% DMSO, 0.01% maltoside in concentration 10 μ M by using the liquid handling robot Biomek 2000. The oligos were printed in duplicate onto aminosilane glass slides activated with PDITC. Using two-color oligo microarrays, breast cancer samples can be compared by hybridizing each sample with a common reference RNA.

C. RNA extraction

Breast cancer tissues were homogenized by using the homogenizer Dia Max of HEIDOLPH and total RNA was

extracted with Qiazol Lysis reagent (Qiagen) and further purified on Rneasy columns (Qiagen) according to the manufacturer's instructions.

D. Probe preparation

To obtain enough amplified RNA for a microarray experiment, a round of RNA amplification was performed on all samples. To serve as reference in microarray hybridizations, a human universal reference RNA from Stratagene was amplified identically. Reverse transcription was performed in the presence of 10mM each of dATP, dCTP, dGTP, dTTP (Invitrogen), 0.1M dithiothreitol, 5x First strand buffer, and 200U Superscript III (Invitrogen). The second strand synthesis of cDNA was catalyzed by 20U of E.coli DNA polymerase I (New England Biolabs) while 2U of RNase H (Invitrogen) produces the primers for the cDNA synthesis. The cDNA molecules were transcribed by T7 RNA polymerase (Epicentre) in 20 μ l reactions at 42°C for 6 hours. The amplified RNA was purified on Rneasy columns (Qiagen) and quantified in Nanodrop.

E. Probe Labeling and Hybridization

The amplified RNA (10 μ g) was diluted in 2 μ l of 0.5M sodium bicarbonate buffer and 10 μ l DMSO. Three mixes with different combinations of external RNA controls were spiked into RNA samples. External controls are RNA molecules that are synthetically produced by in vitro transcription. NHS ester of Alexa 647 was added to the tissue RNA reaction and Alexa 555 dye (Molecular probes) was added to the reference RNA reaction. Both reactions were incubated at 50°C for 3 hours. The unincorporated dye was removed using Microspin G-50 columns (GE Healthcare). Labeling efficiency and quantity of labeled RNA was determined with the spectrophotometer Nanodrop. The ratio of unlabeled to labeled nucleotides was typically between 20 to 30 bases. Slides were prehybridized in 5xSSC, 0.1% SDS, 1%BSA for 90 min and washed with 5x SSC, 0.1% SDS. The labeled probes Cy3 and Cy5 were combined and diluted in 85.5 μ l hybridization buffer (5x SSC, 0.1% SDS, 50% formamide). 15 μ g fragmented salmon sperm DNA were added to combined samples that were subsequently denatured at 80°C for 5 min. Hybridizations were carried out at 48°C for 16h, followed by washing in : 2x SSC, 0.1% SDS at 42°C twice, 0.1X SSC, 0.1% SDS at room temperature twice, and 0.1X SSC at room temperature three successive runs.

F. Scanning and image processing

Arrays were scanned using a GSI Lumonics ScanArray5000. Data were collected in Cy3 and Cy5 channels and stored as TIFF images. Fluorescent intensities of Cy5 and Cy3 channels on each slide were subjected to spot filtering and normalization. In a microarray study, "normalization" indicates the attempt to identify and remove systematic sources of variation in the measured intensities due to separate reverse transcription and labeling, different scanning parameters etc. In our study the normalization was performed by using the print-tip loess normalization method which is one of the most commonly utilized normalization techniques.

G. Statistical analysis

We applied the Significance Analysis of Microarrays (SAM) statistic [6] to detect the genes that are differentially expressed in ER+ with respect to ER- samples. This method implies a modified t-statistic in order to reveal the significant genes. More specifically, for every gene i , it calculates a metric d_i . The larger this metric, the more significant the gene is.

$$d_i = \frac{r_i}{s_i + s_0} \quad (1)$$

where r_i is the mean difference of each gene's i expressions between the two groups, s_i is the standard deviation of gene's i expression across all samples and s_0 is a small number used to avoid the possibility that the fraction goes to infinity. The interested reader can refer to the SAM manual [5] for further information. The SAM method has been widely used in the past [3] and it is a well recognized method in detecting differentially expressed genes. It is the method the more cited for microarray analysis according to the Science Citation Index. In this method, repeated permutations of the data are used to determine if the expression of any gene is significantly related to ER expression. Thus, expected scores for the significance of the genes are retrieved and compared to their observed scores. The use of permutations gives an allure to this method and is recommended when statisticians want to obtain a truly significant result.

H. Data

For the computational analysis described hereafter we used the R [24] language. Before proceeding to the main analysis, we reduced the size of the data in order to minimize computing power and increase efficiency. Thus, we applied a 2-fold filter to the genes, meaning that we kept only the genes that were more than 2-fold up-down-regulated in the examined tissues with respect to the reference sample. This procedure resulted in 3,653 genes, with which we did our manipulations, from the initial set of 34,000 genes.

SAM analysis requires that its input data are linearly normalized across genes and across samples [5]. We performed print tip loess normalization across the genes expressed in each tumor; Between array normalization was not required as the means of the gene expressions and the standard deviations between the samples (arrays) were almost equal. It should be assured that the 3,653 gene expressions in each sample have median 0 according to [5]. The data were thus scaled to median=0 and we ensured that they follow the normal distribution depicted in Figure 1.

To exhaustively search the research literature for genes and biological processes we used Biolab Experiment Assistant (BEA) [22], [23] and DAVID tools [25].

III. RESULTS

After performing the SAM analysis for our two-class unpaired data, we set the δ parameter according to the allowed False Discovery Rate (FDR). The FDR was selected in a way that it was as small as possible (ideally 0) but corresponding to a descent number of genes. It has been

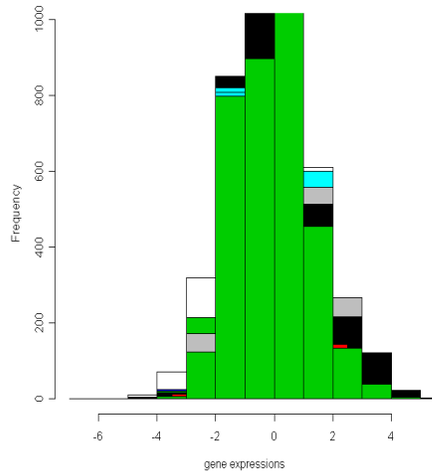


Fig. 1. Superimposed histograms of the distributions of the 3653 gene expressions in each sample. The distributions are normal and that allows us to apply our computational techniques

declared that an FDR of up to 10% – 20% is allowed [7]. We chose a δ of approximately 0.52 which corresponds to an FDR of 9%. In this case we retrieved 112 statistically differentially expressed significant genes, 51 of which are ER+ indicators and 61 of which are ER- indicators.

We were interested in comparing the retrieved results with other methods, to check the consistency of the methods. One of the secondary methods that we applied is the *Rank Products* method [9]. By taking a cutoff at the first 60 more significant ER- correlated genes and at the first 60 more significant ER+ correlated genes, we tried to detect the genes which overlap with our signature. We detected 30 common genes in the ER+ group and 31 common genes in the ER-group. Another method that we applied was the *Pearson Correlation Coefficient* with ER status. This method was adapted from Van't Veer et. al. [1]. To be more specific, we calculated the correlation of each gene's expression across experiments with ER status, which is defined as a vector of 0 and 1 values (0 corresponds to ER- and 1 to ER+ tissues). We randomly put a correlation coefficient cutoff at +0.5 and at -0.5. The genes whose correlation coefficient was > 0.5 are highly expressed in ER+ samples and the ones whose correlation coefficient was $< (-0.5)$ are highly expressed in ER- samples. In the first case we found 55 genes and in the second 149 genes. 42 of our ER+ correlated genes and 55 of our ER- correlated genes were found in these lists. The above findings are very encouraging; they indicate that the differences in gene expressions reflect real biological significance rather than just statistical significance.

A. Clustering

The heatmap of the gene expression levels of the retrieved significant genes is given in Figure 2. Hierarchical clustering using *euclidean* distance and *complete* linkage is applied to both genes and tumors. Almost the same clustering is retrieved when other distances and other types of linkage are

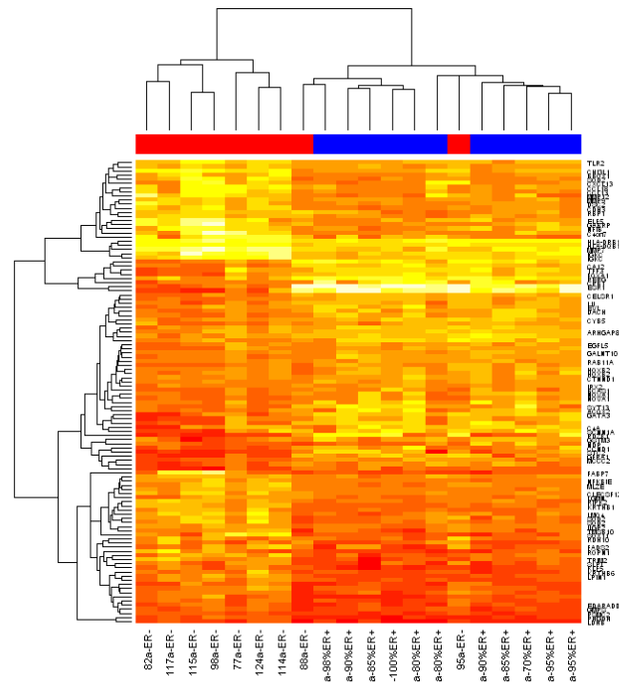


Fig. 2. Expression levels of the significant genes across the 20 samples. Yellow indicates overexpression and red underexpression. The columns correspond to the breast cancer samples. The real status of the samples is given in a color scale: blue for ER+, red for ER-.

used. The genes are put in clusters with expression levels indicative of the sample category. The real status of the samples is given in a color scale: blue for ER+, red for ER-. As it can be noticed there exist two cases where ER- samples 88a and 95a are put in the wrong cluster; thus they seem to be ER+ although they are declared as ER-. By recurring to the raw data, these two samples have indeed values that are closer to the ones observed in ER+ than to the ones observed in ER- samples. As far as the clustering of genes is concerned, three main clusters can be observed; two with high expression in ER- samples and low in ER+, and one cluster with the opposite pattern of expression. In one of the two high in ER- clusters, the high gene expressions are much higher than in the other.

B. Classification

In this section we tried to classify our tumors into ER+ and ER- category. Thus we used a method very similar to the one reported by Van't Veer et al. [1]. We took one sample out and measured its correlation coefficient (Pearson correlation) with the average good and poor prognosis expression levels of the remaining samples. Only the expression levels of the 112 significant genes across the samples were used. We repeated this procedure 20 times, so as to cover all the samples. This method's results regarding the two also mentioned before cases (88a and 95a) were discrepant when correlated to immunohistological data produced by the Pathologists. Because of the clustering and classification results regarding these two samples the slides were reviewed. Indeed, while the infiltrating component of the carcinoma was negative in

these cases, in the representative tissue sections an ER+ in situ component (routinely not reported by Pathologists) was present (in a percentage of about 1%), which was detected by our molecular method, most evidently.

C. A new signature

Taking the above results into account, we decided to exclude from our analysis the two samples for which the errors were reported. and extract a new signature using the remaining 18 samples. Out of these samples, 11 are ER+ and 7 ER-. The method used was the Significance Analysis of Microarrays (SAM) and the data were confirmed to be correctly normalized also in this case. Using this set of samples, we extracted 97 significant genes, 61 of which were significantly higher expressed in the ER+ samples and 36 of which were significantly higher expressed in the ER- samples. A full list of the significant probes and of corresponding gene symbols (in the annotated cases) is given in Supplementary Information 1. Tables I and II represent significant probes and genes along with information on biological pathways and published involvement in breast cancer. These tables were populated with the use of *R* [24] functions and the DAVID tool [25]. More genes that are connected with Gene Ontology (GO) functions can be found in the supplementary files *ER positive* and *ER negative* genes. The clustering of the 18 tumors based only on the 97 significant genes is given in Figure 3. In this case too, *euclidean distance* and *complete linkage* were used for the clustering procedure. As it can be noticed, this time all 18 samples were put in the correct cluster. In order to better validate our signature, we also applied a classification method similar to the one we applied when the 20 samples took part in the analysis. All 18 samples were assigned to the correct category. We were also very interested in examining if these 97 significant genes can assign to the correct cluster all the samples which we analyzed using microarrays (26 samples). In this case the two problematic samples, 88a and 95a, were labeled as ER+. By applying the same clustering method as above we retrieved the heatmap of Figure 5. As it can be noticed, there exist two samples, 77a and 114a which, although they are ER-, they are put in the ER+ cluster. Nevertheless, two ER+ clusters exist; one containing the highly expressed (%) ER+ samples and one containing the less expressed samples. The reported ER- samples were assigned to this second one cluster which justifies in a way this result. Of 97 genes, 19 overlap with the signature found by Veer et.al. [1] and 33 with the signature by West et.al. [4]. This observation is very encouraging; it verifies that there exist some genes that can indeed be indicative of ER status. A full list of the genes found to be in common between the datasets is provided with Supplementary Information 2.

D. Optimal signature

Scientists around the world are mostly interested in minimizing the false positive rate (fpr) using as less genes as possible. Thus, they try to find a sub-signature of the initial gene signature which has the same, or even better result,

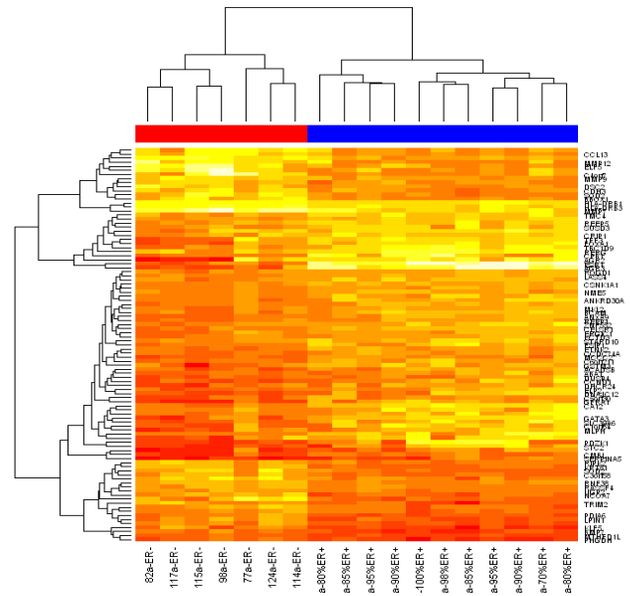


Fig. 3. Expression levels of the significant genes across the 18 samples. Yellow indicates overexpression and red underexpression. The columns correspond to the breast cancer samples. The real status of the samples is given in a color scale: blue for ER+, red for ER-.

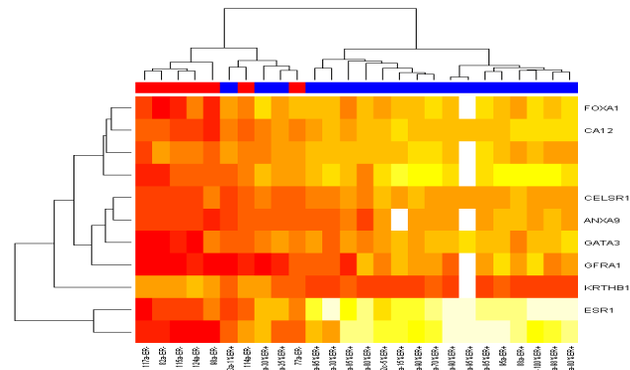


Fig. 4. Heatmap of the 11 most significant genes across the samples. Yellow color indicates overexpression and red underexpression. There exists 1 gene that is up-regulated in ER- tumors and 10 genes that are upregulated in ER+ tumors. The clustering of tumors and genes is made using euclidean distance and complete linkage.

compared to the original one. One of the most known papers that performs this procedure is the paper of Van't Veer et.al [1]. Taking into account this paper we decided to follow the analyzed procedure:

- Rank the retrieved genes in decreasing order according to the correlation of their score with the ER status of the patients
- Take one by one the genes, starting from the top of the ordered list, and classify the tumors based on their expression profiles
- Define the cutoff point, meaning the minimum number of genes having the minimum error

In our case, we ranked the genes and started classifying the tumors using a signature starting from 2 genes and ending at 97 genes, adding one gene at each step. The number of misclassifications was high for 2 genes and continued

TABLE I
ER- indicator genes

Probe id	Gene name	Pathway	BC association
H20000442	MMP12		
H20000574	MMP7	Signal transduction (wnt path)	Associated with poor prognosis, wound healing and metastasis)
H200013790	MMP9		Tumor invasion and angiogenesis in BC and other cancers
H200015680	DSC2	Cell communication	
H300022173	KRTHB1	Cell communication	
H20000695	CDH3	Signaling molecules	Tumor aggressiveness in BC LOH events of Chr16p in BC
H300022893	HLA-DRB1	Signaling molecules, immune system	
H300021886	HLA-DRB3	Signaling molecules, immune system	
H200001995	CCL13	Signaling molecules, immune system	
H200005719	GABRP	Signaling molecules: Neuroactive ligand-receptor interaction	Down regulated in BC, Index of tumor progression, prognostic marker
H300004703	CCL18	Signaling molecules: Cytokine-cytokine receptor interaction	
H200007916	BMP7	Signaling molecules, signal transduction Expressed in BC, may associated with bone metastasis	Expressed in various breast cancer cell lines
H200007119	KLF5		Suppresses tumor cell growth in breast cancer
H200006022	CHI3L1		High serum levels of YKL-40 associated with poor prognosis
H200004198	SOX11		Play a role in tumorigenesis
H200009652	RASSF4		Potential tumor suppressor. May promote apoptosis and cell cycle arrest
H300021244	UGP2	Carbohydrate metabolism,	Forms UDP-glycose which in mammary lactating gland is converted to Udp-galactose and lactose
H200004673	MTHFD1L	Carbohydrate metabolism, metabolism of cofactors and vitamins	
H300006924	PHGDH	Aminoacid metabolism	
H300002199	BBOX1	Aminoacid metabolism (lysine degradation)	

TABLE II
ER+ indicator genes

Probe ID	Gene name	Pathway	BC association
H200000435	ESR1		Stimulation of growth of breast cancer. Response to endocrine therapy
H300000645	NME5		Confers protection from cell death by Bax and alters the cellular levels of several antioxidant enzymes including Gpx5.
H200007883	GATA3		Highly associated with ER & PgR, tumor grade. Involved in growth control & maintenance of differential state in epithelial cells
H300003818	FOXA1		Decreased expression in BC. Mediated ER in BC cells
H200003045	RERG		Loss may contribute to tumorigenesis in breast. Decreased in BC with poor prognosis
H200010467	AGR2		Associated with ER+ BC. Interacts with metastasis genes Potential therapeutic target and molecular marker in prostate cancer
H200014049	STC2		Expression induced by estrogen, altered in BC
H200006989	CCND1	Cell growth and death	Regulated positively by Rb. Mutations, amplification and overexpression of this gene, are observed frequently in a variety of tumors and may contribute to tumorigenesis. Better outcome
H200006652	BCL2	Upregulated in response of human prolactin treatment in BC cancer cell lines.	Expressed in BC, inverse correlation with cytological grade
H300002542	NAT1/NAT2	Caffeine metabolism, drug metabolism	Polymorphisms associated with BC risk
H200006150	DHCR24	Protects cells from oxidative stress by reducing caspase 3	Activity during apoptosis induced by oxidative stress
H200000512	GSTM3	Aminoacid metabolism	
H200017772	ABAT	aminoacid and carbohydrate metabolism	
H200007735	MCCC2	Aminoacid metabolism	
H300004674	ETNK2	Lipid metabolism	
H200006864	ACADSB	Lipid metabolism	
H200001041	CA12	Energy metabolism	
H300015296	CSNK1A1	Signal transduction	Association with BC metastasis
H200000600	DUSP4	Signal transduction	
H200014021	BCAM		Up-regulated following malignant transformation in some cell types. Play a role in epithelial cell cancer
H200006636	SLC39A6		Better outcome in BC. Upregulated by estrogen in BC cell lines.
H200016503	DNAJC12		(DeBessa SA 2006) Correlation with ER in BC
H300003702	PDZK1		May play a role in the cellular mechanisms associated with multidrug resistance through its interaction with ABC2 and PDZK1IP1.
H200019227	ANKRD30A		NY-BR-1 is a differentiation antigen present in BC Possible antigenic target for antibody treatment
H200020432	CMBL	Xenobiotics biodegradation	
H200006282	SERPINA5	Immune system	Positive prognostic factor (suppression of tumor invasion)

- [4] M. West, C. Blanchette, H. Dressman, E. Huang, S. Ishida, R. Spang, H. Zuzan, J. A. Olson, Jr., J. R. Marks and J. R. Nevins, "Predicting the clinical status of human breast cancer by using gene expression profiles", *PNAS*, vol. 98, no. 20, pp 11462-11467 (2001)
- [5] G. Chu, B. Narasimhan, R. Tibshirani and V. Tusher, "SAM Significance Analysis of Microarrays Users guide and technical document"
- [6] V. Tusher, R. Tibshirani, and G. Chu., "Significance analysis of microarrays applied to transcriptional responses to ionizing radiation.", *Proc. Natl. Acad. Sci. USA.*, vol. 98, pp 5116-5121 (2001)
- [7] S. G. Baker and B. S. Kramer, "Using microarrays to study the microenvironment in tumor biology: The crucial role of statistics.", *Seminars in Cancer Biology* (2008), doi:10.1016/j.semcancer.2008.03.001
- [8] G. Potamias, A. Analyti, D. Kafetzopoulos, M. Kafousi, T. Margaritis, D. Plexousakis, P. Poirazi, M. Reczko, I.G. Tollis, M. E. Sanidas, E. Stathopoulos, M. Tsiknakis, S. Vassilaros, "Breast Cancer and Biomedical Informatics: The PrognoChip Project", *IMACS 2005: Computer Science and Artificial Intelligence - Bioinformatics session*, Paris, France, July 11-15 (2005)
- [9] R. Breitling, P. Armengaud, A. Amtmann. and P. Herzyk, "Rank products: a simple, yet powerful, new method to detect differentially regulated genes in replicated microarray experiments", *FEBS Letters*, vol. 573, pp 83-92 (2004)
- [10] H. P. Koeffler, C. W. Miller, B. Y. Karlan, I. Wolf, S. Bose and E. A. Williamson, "FOXA1: Growth inhibitor and a favorable prognostic factor in human breast cancer.", *Int J Cancer*, vol. 120 , pp 1013-22 (2007)
- [11] M. Brown and J. S. Carroll, "Estrogen receptor target gene: an evolving concept.", *Mol Endocrinol*, vol. 20 , pp 1707-14 (2006)
- [12] J. Schneider, M. Ruschhaupt, A. Buneß, M. Asslaber, P. Regitnig, K. Zatloukal, W. Schippinger, F. Ploner, A. Poustka and H. Sultmann, "Identification and meta-analysis of a small gene expression signature for the diagnosis of estrogen receptor status in invasive ductal breast cancer", *Int. J. Cancer*, vol. 119, pp 2974-2979 (2006)
- [13] T. Bouras, M. C. Southey, A. C. Chang, R. R. Reddel, D. Willhite, R. Glynn, M. A. Henderson, J. E. Armes and D. J. Venter, "Stanniocalcin 2 Is an Estrogen-responsive Gene Coexpressed with the Estrogen Receptor in Human Breast Cancer", vol. *Cancer Research*, vol. 62, pp 1289-1295 (2002)
- [14] N.M. Malara, A. Leottab, A. Sidotia, S. Liob, R. D'Angelo, B. Caparellob, F. Munaoc, F. Pinoa and A. Amato, "Ageing, hormonal behaviour and cyclin D1 in ductal breast carcinomas", *The Breast*, vol. 15, Issue 1, pp 81-89 (2006)
- [15] S. Tozlu, I. Girault, S. Vacher, J. Vendrell, C. Andrieu, F. Spyrtos, P. Cohen, R. Lidereau and I. Bieche, "Identification of novel genes that co-cluster with estrogen receptor alpha in breast tumor biopsy specimens, using a largescale real-time reverse transcription-PCR approach", *Endocrine-Related Cancer*, vol. 13, pp 1109-1120 (2006)
- [16] M. Lacroix, "Significance, detection and markers of disseminated breast cancer cells", *Endocrine-Related Cancer*, vol. 13, pp 1033-1067 (2006)
- [17] A. J. Minn, G. P. Gupta, P. M. Siegel, P. D. Bos, W. Shu, D. D. Giri, A. Viale, A. B. Olshen, W. L. Gerald and J. Massague, "Genes that mediate breast cancer metastasis to lung", *Nature*, vol. 436, pp 518-524 (2005)
- [18] S. Khandpur, R. Bamezai, B. S. N. Reddy and N. K. Bairwa, "A study of phenotypic correlation with the genotypic status of HTM regions of KRTHB6 and KRTHB1 genes in monilethrix families of Indian origin.", *Ann Genet*, vol. 47, pp 77-84 (2004)
- [19] S. Mochizuki, M. Shimoda, T. Shiomi, Y. Fujii and Y. Okada, "ADAM28 is activated by MMP-7 (matrilysin-1) and cleaves insulin-like growth factor binding protein-3", *Biochemical and Biophysical Research Associations*, vol. 315, Issue 1, pp 79-84 (2004)
- [20] P. Neven, R. Paridaens, H. Wildiers, A. Smeets, W. Hendrickx, M. Drijkoningen, J. Decock, "Matrix metalloproteinase expression patterns in luminal A type breast carcinomas.", *Dis Markers*, vol. 23, pp 189-96 (2007)
- [21] P. M. McGowan and M. J. Duffy, "Matrix metalloproteinase expression and outcome in patients with breast cancer: analysis of a published database", *Annals of Oncology*, doi:10.1093/annonc/mdn180 (2008)
- [22] URL: <http://www.biovista.com/bea>
- [23] D. Mastellos, C. Andronis, A. Persidis and J. D. Lambris, "Novel biological networks modulated by complement", *Clin Immunol.*, vol 115, issue 3, pp 225-35 (2005)
- [24] URL: <http://www.r-project.org/>
- [25] URL: <http://david.abcc.ncifcrf.gov/>
- [26] C. R. Acharya, D. S. Hsu, C. K. Anders, A. Anguiano, K. H. Salter, K. S. Walters, R. C. Redman, S. A. Tuchman, C. A. Moylan, S. Mukherjee, W. T. Barry, H. K. Dressman, G. S. Ginsburg, K. P. Marcom, K. S. Garman, G. H. Lyman, J. R. Nevins and A. Potti, "Gene Expression Signatures, Clinicopathological Features, and Individualized Therapy in Breast Cancer", *Jama*, vol. 299, No. 13, pp 1574-1587 (2008)
- [27] H. Nakshatri and S. Badve, "FOXA1 as a therapeutic target for breast cancer", *Expert Opin Ther Targets*, vol. 11, pp 507-14 (2007)
- [28] L. Ein-Dor, I. Kela, G. Getz, D. Givol and E. Domany, "Outcome signature genes in breast cancer: is there a unique set", *Bioinformatics*, vol. 21, pp 171-178 (2005)
- [29] G. M. Callagy, M.J. Webber, P.D Pharoah and C. Caldas, "Meta-analysis confirms BCL2 is an independent prognostic marker in breast cancer", *BMC Cancer*, vol. 8, issue 153 (2008)