

# Gene expression is highly correlated on the chromosome level in urinary bladder cancer

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**Keywords:** bladder cancer, chromosome correlation maps, molecular networks, mathematical modeling, data simulations

**Objective:** Chromosome correlation maps display correlations between gene expression patterns on the same chromosome. Our goal was to map the genes on chromosome regions and to identify correlations through their location on chromosome regions.

**Results:** The top deregulated molecules among 129 bladder cancer samples were implicated in the PI3K/AKT signaling, cell cycle, Myc-mediated apoptosis signaling and ERK5 signaling pathways. Their most prominent molecular and cellular functions were related to cell cycle, cell death, gene expression, molecular transport and cellular growth and proliferation. Chromosome correlation maps allowed us to detect significantly co-expressed genes along the chromosomes. We identified strong correlations among tumors of T $\alpha$ -grade 1, as well as for those of T $\alpha$ -grade 2, in chromosomes 1, 2, 3, 7, 12 and 19. Chromosomal domains of gene co-expression were revealed for the normal tissues, as well. The expression data were further simulated, exhibiting an excellent fit ( $0.7 < R^2 < 0.9$ ). The simulations revealed that along the different samples, genes on same chromosomes are expressed in a similar manner.

**Materials and Methods:** Following microarray analysis we used Ingenuity Pathway Analysis (IPA) to construct gene networks of the co-deregulated genes in bladder cancer. Chromosome mapping, mathematical modeling and data simulations were performed using the WebGestalt and Matlab<sup>®</sup> softwares.

**Conclusions:** Gene expression is highly correlated on the chromosome level. Chromosome correlation maps of gene expression signatures can provide further information on gene regulatory mechanisms. Gene expression data can be simulated using polynomial functions.

## Introduction

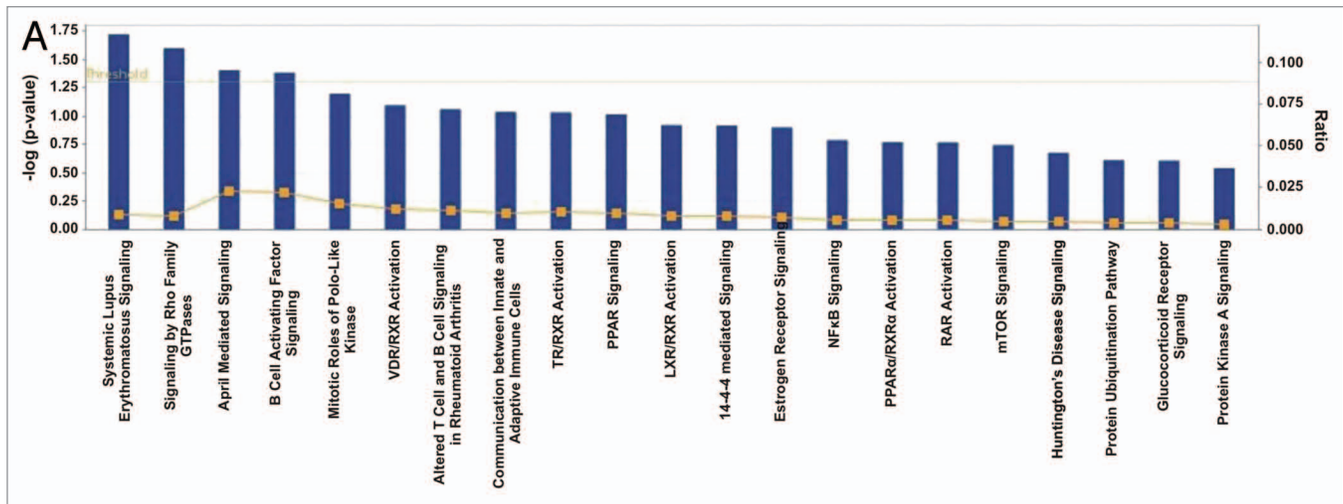
Microarray analyses have been applied in the past to examine the differentially expressed (DE) genes in bladder cancer.<sup>1-7</sup> However, these findings are not always reproducible, mainly due to improper analysis/validation, insufficient control of false positives, inadequate reporting of methods and small sample size relative to large numbers of potential predictors. We have previously shown that the combination of information from multiple studies can increase the reliability and recognizability of the results, and can lead us to the focus of genes that play a role in the formation of bladder cancer, irrespective of the stage and/or grade of the tumor.

Such high-throughput analyses can provide a system-scale overview of how genes interact with each other in a network context. This network is known as a gene regulatory network and can be defined as a mixed graph over a set of nodes (corresponding to genes or gene activities) with directed or undirected edges (representing causal interactions or associations between gene

activities).<sup>8,9</sup> As most biological processes arise from the complex interactions among multiple gene products, information about how genes function together can improve our understanding of the underlying biological mechanisms.

Chromosome correlation maps display correlations between the expression patterns of genes on the same chromosome and are considered of major importance in the understanding of gene expression regulation.<sup>10,11</sup> Considering that gene expression represents just a “snap shot” of the state-space of the otherwise dynamic behavior of bladder cancer, it would be of major interest to investigate the expression patterns by examining the chromosomal-based gene expression. Chromosomal gene expression is expected to be highly coordinated. However, it has not yet been elucidated based on chromosome correlation whether gene expression among same chromosomes from different samples is governed by similar patterns. If such a common mechanism of gene expression exists, we do not know whether it is of linear or nonlinear nature.

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**Figure 1A.** Ingenuity analysis of the top pathways affected in differentially expressed genes among 10 bladder cancer and five normal tissue samples (Cohort A). Y-axis is an inverse indication of p-value or significance (A). Gene networks involved in “cell-to-cell signaling and interaction, cellular assembly and organization, cellular function and maintenance” (B), and “cell signaling, molecular transport and nucleic metabolism” (C), generated by IPA for differentially expressed genes between bladder cancer and normal tissue. The selected scoring method was Fisher’s exact test p-value. The threshold value was set at  $p = 0.05$ . Red symbols are assigned for upregulated and green for downregulated genes. Node shape corresponds to the functional role of molecules as shown in the legend. Direct or indirect interactions are shown by complete or dashed lines.

Our goal was to construct networks of the co-deregulated (co-DE) genes in bladder cancer, to unveil the gene correlations among tumor samples and to describe these correlations mathematically in order to perform predictions of gene expression based on the chromosomes.

## Results

**Networks and canonical pathways.** *Networks and canonical pathways for the DE genes revealed in Cohort A.* We performed Ingenuity Pathway Analysis (IPA) for 831 co-DE genes among the ten bladder cancers and the five normal tissue samples, in Cohort A. The top canonical pathways were as follows: (1) Systemic Lupus Erythematosus signaling ( $p = 1.9E-02$ ; ratio = 0.009), containing the genes LSM5 and SNRPG (both upregulated); (2) signaling by Rho family GTPases ( $p = 2.52E-02$ ; ratio = 0.008), containing the genes DES and GFAP (both downregulated); (3) April-mediated signaling ( $p = 3.94E-02$ ; ratio = 0.023), containing the gene TNFRSF17 (downregulated); (4) B cell-activating factor signaling ( $p = 4.15E-02$ ; ratio = 0.022), containing the gene TNFRSF17 (downregulated); and (5) mitotic roles of Polo-like kinase ( $p = 6.36E-02$ ; ratio = 0.015), containing the gene ANAPC11 (upregulated). Two major gene networks were constructed, with the following associated functions: (1) cell-to-cell signaling and interaction, cellular assembly and organization, cellular function and maintenance (score = 36) and (2) cell signaling, molecular transport and nucleic metabolism (score = 11) (Fig. 1A–C).

The top molecular and cellular functions were: (1) cell death ( $p = 1.06E-03 - 2.20E-02$ ; three molecules: LTF, TNFRSF17 and GFAP, all downregulated); (2) cell morphology ( $p = 1.06E-03 - 2.09E-02$ ; three molecules: LTF, DES and GFAP,

downregulated; and NCOR1, upregulated); (3) cellular assembly and organization ( $p = 1.06E-03 - 4.25E-02$ ; eight molecules: CAMSAP3 and NCOR1, upregulated; and FHL1, DES, CAP2, GFAP, LTF and TNFRSF17, downregulated); (4) cellular compromise ( $p = 1.06E-03 - 2.11E-03$ ; DES, downregulated) and (5) cellular function and maintenance ( $p = 1.06E-03 - 4.25E-02$ ; seven molecules: CAMSAP3, upregulated; and GFAP, CAP2, DES, LTF, PEX5L and FHL1, downregulated).

The top upregulated molecules were: C7orf68, ANAPC11, NCOR1, LSM5, SNRPG, GDAP2, MAPKAP1, TOX4 and CAMSAP3, whereas the top downregulated molecules were TAC3, LTF, ODZ2, FHL1, TNFRSF17, DES, ANKRD29, GPSM1, QRICH2 and GFAP (Table 1). The top upstream regulators were as follows: KDM4A (targeting FHL1), RNF112 (targeting TAC3), ALX3 (targeting GFAP), SOX11 (targeting C7orf68) and OLIG2 (targeting GFAP) ( $p = 1.06E-03 - 5.27E-03$ ).

*Networks and canonical pathways for the DE genes revealed in Cohort B.* Similarly, we performed IPA for the 458 co-DE genes among the 129 bladder cancers and the 17 normal tissue samples in Cohort B. The top canonical pathways were as follows: (1) PI3K/AKT signaling ( $p = 3.7E-04$ ; ratio = 0.021), containing the genes CTNNB1, YWHAB and YWHAE (all upregulated); (2) cell cycle: G<sub>2</sub>/M DNA damage checkpoint regulation ( $p = 1.25E-03$ ; ratio = 0.041), containing the genes, YWHAB and YWHAE (both upregulated); (3) Myc-mediated apoptosis signaling ( $p = 2.24E-03$ ; ratio = 0.033), containing the genes YWHAB and YWHAE (both upregulated); (4) ERK5 signaling ( $p = 2.47E-03$ ; ratio = 0.031), containing the genes YWHAB and YWHAE (both upregulated); and (5) basal cell carcinoma signaling ( $p = 3.14E-03$ ; ratio = 0.027), containing the genes CTNNB1 (upregulated) and WNT10B (downregulated) (Fig. 2A–C).

The top molecular and cellular functions were: (1) cell cycle ( $p = 1.66E-07 - 1.54E-02$ ; 11 molecules: CDC20, CTNNB1,

MARCKS, PCNA, PSEN1, KRT7, YWHAE, YWHAB, upregulated; and CD40LG, TPM2, WNT10B, downregulated); (2) cell death ( $p = 1.79E-05 - 1.95E-02$ ; 12 molecules: BCLAF1, CDC20, CTNNB1, PCNA, CTNNB1, PSEN1, YWHAE, YWHAB, upregulated; and ACTC1, CD40LG, GLP1R, downregulated); (3) gene expression ( $p = 2.65E-05 - 1.54E-02$ ; 12 molecules: BCLAF1, CTNNB1, KRT7, PCNA, PSEN1, YWHAB, YWHAE, BCLAF1, ZFP, upregulated; and CD40LG, GLP1R, TAGLN, WNT10B, downregulated); (4) molecular transport ( $p = 4.18E-05 - 1.55E-02$ ; seven molecules: PSEN1, YWHAB, YWHAE, PCNA, MARCKS, upregulated; and CCKAR, GLP1R, downregulated); and (5) cellular growth and proliferation ( $p = 1.83E-04 - 1.90E-02$ ; nine molecules: CTNNB1, PSEN1, CDC20 and PCNA, upregulated; and CD40LG, TPM2, GLP1R, WNT10B, downregulated).

The top upregulated molecules were KRT7, APOBEC3B, CDC20, MARCKS, BCLAF1, ZFP36L2, YWHAE, PSEN1, CTNNB1 and YWHAB, whereas the top downregulated molecules were ACTC1, TPM2, TAGLN, GLP1R, SPARCL1, MFAP4, CDC40LG, WNT10B, CCKAR and HES1 (Table 2). The top transcription factors were as follows: MKL1 (targeting ACTC1, TAGLN and TPM2), KLF5 (targeting ACTC1, TAGLN and TPM2), SMAD3 (targeting GFAP), HTT (targeting CTNNB1, PCNA, TAGLN and TPM2) and FOXA1 (targeting ACTC1, TAGLN and TPM2) ( $p = 1.66E-06 - 1.06E-04$ ).

**Analysis of the co-deregulated genes in Cohort B. Comparison between all tumor and control samples.** We did not detect any correlation between tumor and control samples. Interestingly, on chromosome 1, the mean value of the DE genes was positive, indicating that they are all upregulated. *CDC20*, whose upregulation in bladder cancer has previously been reported,<sup>7</sup> appeared to be the most interesting. Moreover, *GPREL1* on chromosome 4 and *HCCS* on chromosome X were the most active genes among all tumor samples (Fig. S1). Similarly, we compared all the control and tumor samples. Once again, *CDC20* appeared to be

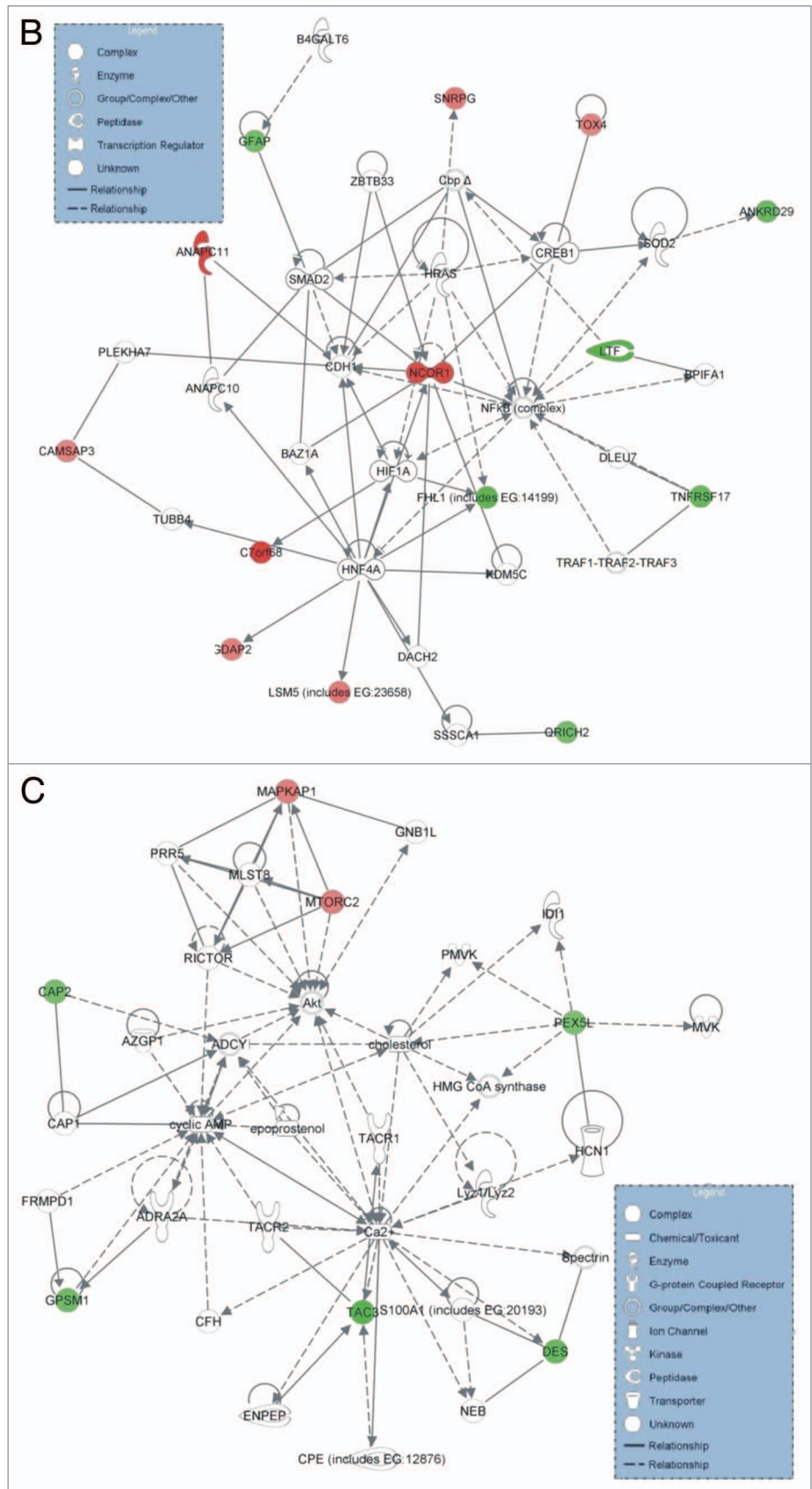


Figure 1B and C. For figure legend, see page 2.

**Table 1.** Top deregulated molecules in cohort A

Upregulated molecules in bladder cancer vs. normal tissue	Exp. value (fold change)
C7orf68	2.633
ANAPC11	2.562
NCOR1	2.433
LSM5 (includes EG:23658)	1.720
SNRPG	1.705
GDAP2	1.635
MAPKAP1	1.624
TOX4	1.580
CAMSAP3	1.570
EPAS1	0.489
Downregulated molecules in bladder cancer vs. normal tissue	Exp. value (fold change)
TAC3	-2.518
LTF	-2.464
ODZ2	-2.438
FHL1 (includes EG:14199)	-2.315
TNFRSF17	-2.065
DES	-2.015
ANKRD29	-1.937
GPSM1	-1.789
QRICH2	-1.739
GFAP	-1.737

10 bladder cancer and five normal tissue samples, as identified by IPA.

upregulated in bladder cancer, thus strengthening its significant implication in the disease. An interesting gene expression pattern was revealed on chromosomes 11, 13, 18, 21 and 22 (Fig. S2). All genes were upregulated on average, indicating that for these chromosomes, their respective genes are the most active in all tumor samples examined.

*Comparison between T $\alpha$ -grade 1 tumors and control samples.* Chromosome correlation maps for T $\alpha$ -grade 1 tumors revealed co-expressed gene patterns along various chromosomes (Fig. 3). Since correlation does not necessarily mean causation, we further searched for possible ways that would describe these patterns of expression. Indeed, these patterns could be described with third degree polynomials, and all simulations manifested an excellent fitting ( $R^2 > 0.99$ ). This was a hint that in this tumor sample, gene expression is highly conserved and related to the nature of the tumor, irrespective of the genes. It was also interesting as the DE genes were randomly selected, since they were produced through a filtering procedure from a t-test. The simulation showed that among different samples, genes on the same chromosome are expressed in a similar manner. Simulation results for all chromosomes are presented in Figure 4.

*Comparison between T $\alpha$ -grade 2 tumors and control samples.* There was a mixture of correlation profiles in T $\alpha$ -grade 2 tumors. In the case of T $\alpha$ -grade 1 tumors, we observed a plethora of correlations with  $\rho > 0.9$  and  $p < 0.05$ . Due to the small sample

**Table 2.** Top deregulated molecules, in cohort B

Upregulated molecules in bladder cancer vs. normal tissue	Exp. value (fold change)
KRT7	2.267
APOBEC3B	2.030
CDC20	2.024
MARCKS	1.882
BCLAF1	1.775
ZFP36L2	1.768
YWHAE	1.655
PSEN1	1.647
CTNNB1	1.577
YWHAB	1.552
Downregulated molecules in bladder cancer vs. normal tissue	Exp. value (fold change)
ACTC1	-3.188
TPM2	-2.694
TAGLN	-2.504
GLP1R	-2.248
SPARCL1	-1.914
MFAP4	-1.755
CD40LG	-1.721
WNT10B	-1.504
CCKAR	-1.502
HES1 (includes EG:15205)	-0.776

129 bladder cancer and 17 normal tissue samples, as identified by IPA

number ( $n = 3$ ), we assumed that fitting, simulation and correlations were close to the ideal. Therefore, in the case of the T $\alpha$ -grade 2 group, chromosome correlation maps were also built in order to allow the visualization of co-expressed genes along the chromosomes (Fig. 5). Simulating gene expression with respect to chromosome location did not give straightforward results, as in the case of the T $\alpha$ -grade 1 tumor group. Polynomial approximations were used, and the  $R^2$  values ranged between 0.7 and 0.91. However, it appeared that they could be simulated with polynomials despite the difficulty to find the most suitable function for the data. This indicates that chromosome-based gene expression is a quasi-linear problem, possibly of a nonlinear nature. The results of the simulations are presented in Figure 6.

*Comparison between T $\alpha$ -grade 3 tumors and control samples.* Similarly, chromosome correlation maps were constructed for T $\alpha$ -grade 3 tumors in order to allow visualization of co-expressed genes along the chromosomes (data not shown due to space limitations).

*Comparison between T1-grade 2 tumors and control samples.* As also expected in the present case, gene expression was highly correlated (Fig. 7). Similarly, simulation of the data showed that they could be fitted easily, as was also the case of the T $\alpha$ -grade 1 samples (data not shown due to space limitations).

*Comparison between T1-grade 3 tumors and control samples.* Chromosome correlation maps for T1-grade 3 tumors did not

exhibit highly co-expressed genes along the chromosomes. The data were not further investigated.

*Comparison between T1-grade 2/3 tumors and control samples.* Chromosome correlation maps for T1-grade 2 and T1-grade 3 tumors also revealed highly co-expressed gene patterns along the chromosomes (data not shown due to space limitations). Similarly, data simulation showed a high ability for fitting, as was also the case of the T $\alpha$ -grade 1 and T1-grade 2 tumor samples (data not shown).

No further correlation could be identified in the chromosome correlation maps for the following tumor comparisons: comparison between T1-grade 2/3/4 tumors and control samples; comparison between group A tumors and control samples; comparison between group B tumors and control; comparison between group C tumors and control samples; and comparison between metastatic tumors and control samples.

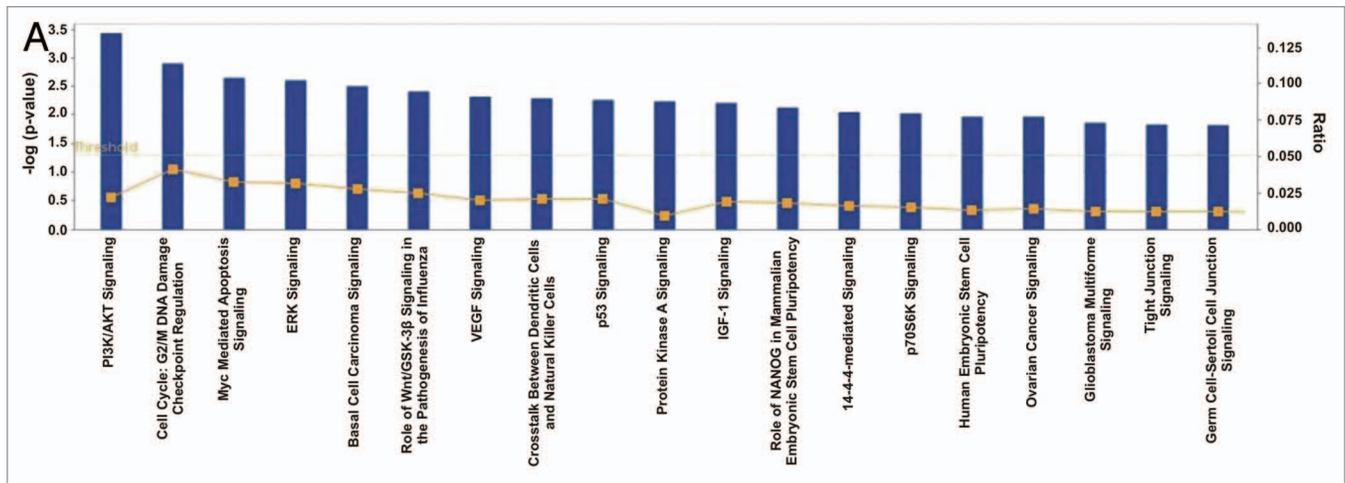
*Comparison among all control samples.* We further investigated the existence of correlations separately among tumor and control samples. Interestingly, the control samples exhibited correlations on certain chromosomes, such as chromosomes 2, 4, 5, 7, 10, 13, 15 and 17 (Fig. S3). Based on the observed correlations, we attempted to fit our data in order to discover patterns of gene expression. If a simulation existed, it should be true for all the samples in specific chromosomes. For this purpose, we calculated the mean values of each sample for all genes and for each chromosome. Surface fittings were selected based on their  $R^2$  value, which should be  $> 0.9$ . This finding was interesting, as it hinted toward an expression pattern in the physiological samples, pointing out that gene expression is indeed highly correlated in cells (Fig. S4).

*Comparison between all tumors and control samples.* The above results suggest a need for a different approach when it comes to analyzing these data. Since gene expression is multidimensional, taking into account that we have 22 chromosomes and two sex chromosomes, the analysis should take place for each one of them. Supposed that we have  $k$  samples with  $j$  genes each and  $c = 22 + 2$  chromosomes, the gene expression  $x$  would be defined as:  $x_{k,j,c}$ . Thus, we need three variables in order to describe the expression of a gene. This leads to the formation of a 3D matrix of the form  $m \times n \times p$ . Yet, such a matrix is a multidimensional structure that cannot be easily visualized. In the present case, in order to visualize gene expression in such a way that would give meaningful interpretation of chromosome-based gene expression, we created this 3D structure. In order to simplify this structure, we isolated 2D matrices of genes vs. samples with constant chromosomes, and genes vs. chromosomes with constant samples. The second problem that we encountered was how to further reduce the complexity of the data and find similarities between gene expression values with respect to chromosomes. 3D visualizations of the initial data were too complex to analyze and to obtain patterns of gene expression. The answer to this question came from the utilization of k-means clustering. K-means classifies gene expressions, based on similarities. Thus, if genes are sorted with respect to chromosomes, then possible patterns could be revealed using the k-means algorithm. In Figure S5, the result of the k-means clustering is presented as implemented for the genes vs.

**Table 3.** Selected GO terms of chromosomal-based gene expression

GO analysis		
ID	Name	p-value (Adj)
<b>chromosome 1</b>		
GO:0060669	embryonic placenta morphogenesis	0.0045
GO:0005839	proteasome core complex	0.0014
GO:0005031	tumor necrosis factor receptor activity	0.0067
<b>chromosome 2</b>		
GO:0009952	anterior/posterior pattern formation	0.0018
GO:0007350	blastoderm segmentation	0.0064
GO:0048468	cell development	0.0071
GO:0048562	embryonic organ morphogenesis	0.0025
GO:0048706	embryonic skeletal system development	0.0062
GO:0005152	interleukin-1 receptor antagonist activity	0.0052
GO:0004918	interleukin-8 receptor activity	0.0052
<b>chromosome 3</b>		
GO:0016493	C-C chemokine receptor activity	4.71E-05
GO:0071425	hemopoietic stem cell proliferation	0.0043
<b>chromosome 4</b>		
GO:0001525	Angiogenesis	0.0052
GO:0008009	chemokine activity	2.99E-07
GO:0008083	growth factor activity	0.0008
GO:0005134	interleukin-2 receptor binding	0.0059
GO:0051781	positive regulation of cell division	0.0001
GO:0045741	positive regulation of epidermal growth factor receptor activity	0.0008
GO:0050729	positive regulation of inflammatory response	0.0027
GO:0045410	positive regulation of interleukin-6 biosynthetic process	0.0003
GO:0045840	positive regulation of mitosis	0.0061
GO:0034105	positive regulation of tissue remodeling	0.0087
GO:0008202	steroid metabolic process	0.0065
<b>chromosome 5</b>		
GO:0042977	activation of JAK2 kinase activity	0.0010
GO:0035240	dopamine binding	0.0020
GO:0008083	growth factor activity	0.0048
GO:0070851	growth factor receptor binding	0.0056
GO:0005138	interleukin-6 receptor binding	0.0068
GO:0004923	leukemia inhibitory factor receptor activity	0.0023
<b>chromosome 6</b>		
GO:0060333	interferon-gamma-mediated signaling pathway	1.78E-09
GO:0046415	urate metabolic process	0.0094
GO:0001570	vasculogenesis	0.0089
<b>chromosome 7</b>		

The terms were considered significant if they obtained an adjusted p-value  $< 0.01$ .



**Figure 2A.** Ingenuity analysis of the top pathways affected in differentially expressed genes among 129 bladder cancer and 17 normal tissue samples (Cohort B). Y-axis is an inverse indication of p-value or significance. (A) Gene networks involved in “cell cycle, gene expression and cell death” (B), and “cell morphology, cellular function and maintenance and cell death” (C), generated by IPA for differentially expressed genes between bladder cancer and normal tissue. The selected scoring method was Fisher’s exact test p-value. The threshold value was set at  $p = 0.05$ . Red symbols are assigned for upregulated and green for downregulated genes. Node shape corresponds to the functional role of molecules as shown in the legend. Direct or indirect interactions are shown by complete or dashed lines.

the samples with constant chromosome location. These results were further fitted using Fourier series (Eqn. 3). The result for chromosome 1 is presented in Figure S6. We further analyzed the data by calculating the confidence intervals at the 95% level, the first and second derivatives, as well as the integral of the fitted data. An interesting observation was that the derivatives of the fitted data manifested oscillatory behavior. In addition, we present the results of our analysis for chromosome X in Figure S7. All remaining chromosomes were analyzed accordingly (data not shown due to space limitations).

Our analysis proceeded with the consideration of chromosomes vs. gene expression. Similarly, we performed k-means clustering analysis (Fig. S8) for the first 10 samples. The remaining 130 samples were clustered accordingly. Simulations of this dimension of the cube were succeeded with Fourier series as mentioned in the “Materials and Methods” section and Equation 3. The result is presented in Figure S9 with indicative diagrams of our analysis. The cubical matrix that was implemented is a very complex structure, which eventually manifests nonlinear dynamics with respect to gene expression.

*Gene ontology (GO) enrichment analysis.* Gene expression was further investigated using GO enrichment. Function distribution along chromosomes is presented in Figure S10. Chromosome 5 presented more functions compared with the other chromosomes. GO enrichment was not proportional to the chromosome size, since the largest chromosome 1 presented 35 significant functions. All the significant gene functions per chromosome are presented in Table 3.

## Discussion

Several studies have focused on the expression profiling of bladder cancer using microarrays.<sup>12-21</sup> In the present study, the bladder

cancer cases were carefully selected in order to obtain at least one pair from each of the following groups: T1-grade 2, T1-grade 3 and T2/T3-grade 3. Furthermore, in our pooled microarray analysis, a wide range of data from publicly available microarray data sets was included, increasing the total number to 129 bladder cancer and 17 control samples. Statistical analysis results were obtained from our previous works,<sup>7,35</sup> and the same collection of genes was used. All conclusions and further analyses were based on the differential gene expression reported. The main difference is that this same data set was further divided according to the chromosomal location of each identified gene.

We performed IPA for two separate cohorts (A and B) and constructed chromosome correlation maps for all bladder tumors in order to visualize co-deregulated genes along the chromosomes. Network analysis manifested two distinct signaling pathways, the glucocorticoid receptor (GR) and NF $\kappa$ B signaling pathway. These pathways are known to be antagonistic, since the second is responsible for the inflammatory response while the first for the anti-inflammatory one.<sup>22</sup> It has been reported that tumors of the urinary tract are not considered to be hormone-dependent, yet it has been shown that nuclear receptors participate in urinary tumor ontogenesis.<sup>23</sup> Also, there are no reports connecting bladder cancer to NF $\kappa$ B. The finding that genes that possibly participate in the progression of the tumor are related to the NF $\kappa$ B signaling pathway is presented for the first time.

The mean value of the DE genes in tumor groups vs. control samples on chromosome 1 was positive, implying that these genes are upregulated. The most interesting case, however, was the appearance of *CDC20*, also previously reported to be significantly overexpressed in bladder cancer.<sup>7</sup> Also, *GPREL1* (chromosome 4) and *HCCS* (chromosome X) were the most active genes among all tumor samples. *CDC20* has been previously reported to act as a potent TP53 target and is a putative therapeutic target gene.<sup>24</sup>

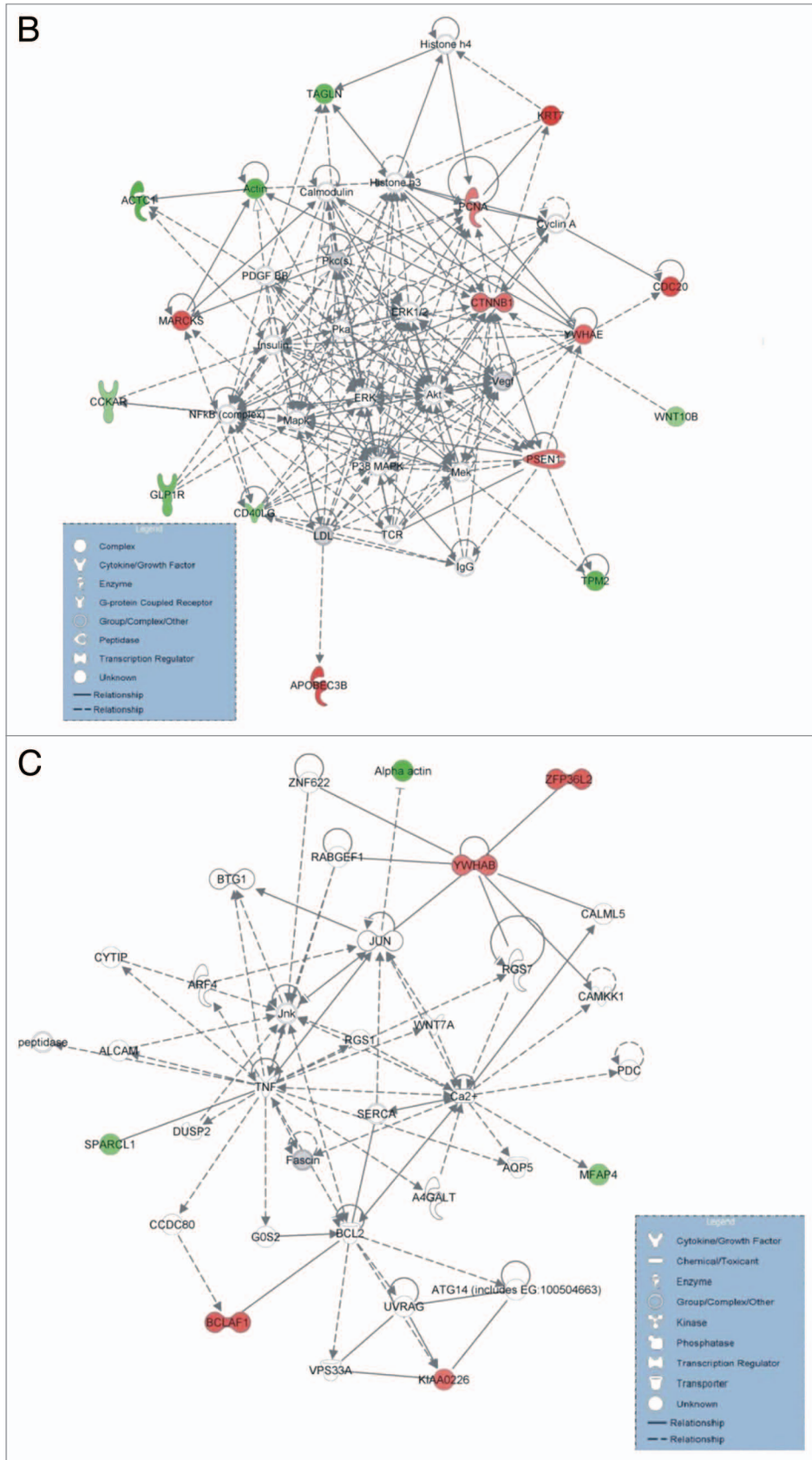
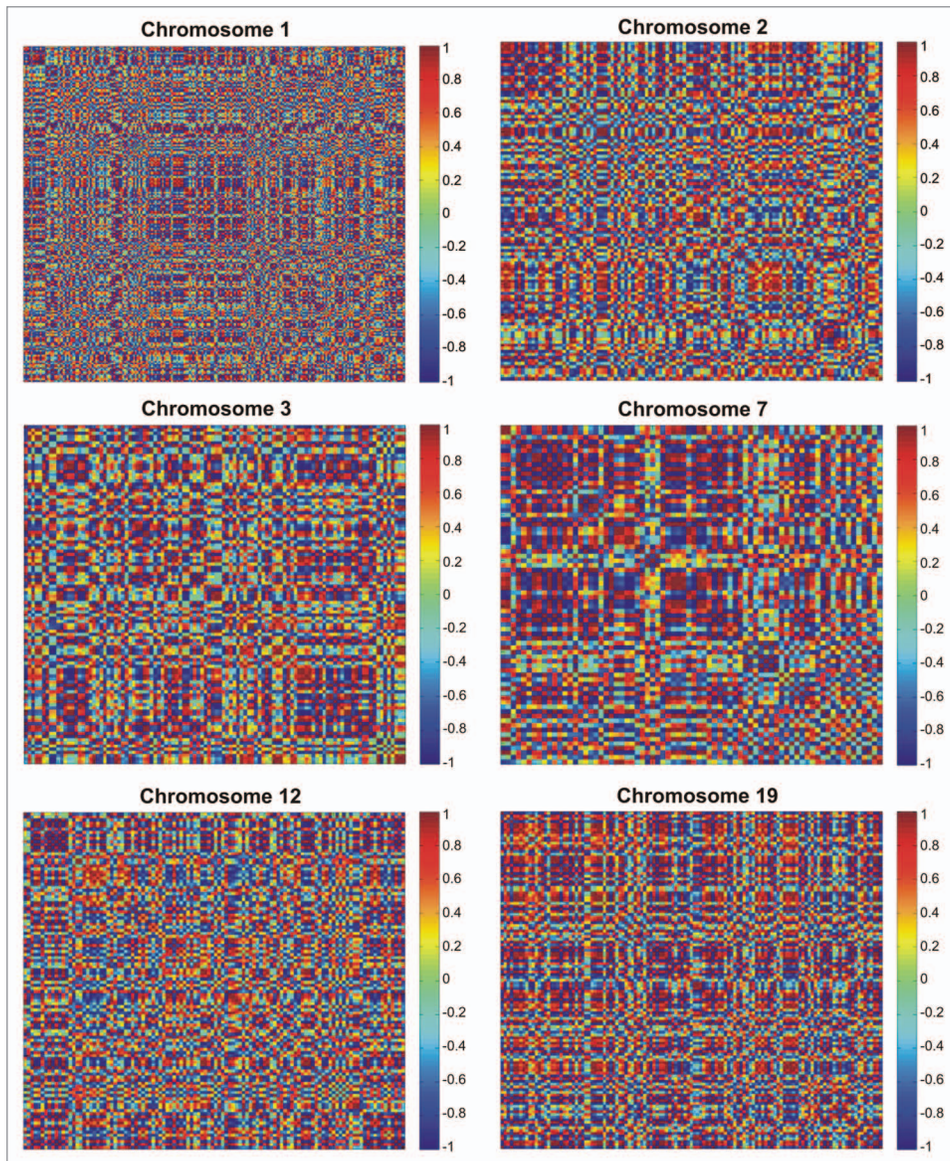


Figure 2B and C. For figure legend, see page 6.



**Figure 3.** Chromosome correlation maps of the DE genes between T $\alpha$ -grade 1 tumors and control samples, on chromosomes 1, 2, 3, 7, 12 and 19. The X and Y axes represent the individual genes that were differentially expressed between control and Ta-grade 1 tumors.

Significantly co-expressed genes along chromosomes 1, 2, 3, 7, 12 and 19 appeared among the tumor subgroups with respect to the control samples. To our knowledge, this is the first report that indicates such gene expression correlations in bladder cancer. Since this was a first indication of gene regulatory mechanism, we wished to ascertain whether gene expression could be simulated. It is known that the existence of correlation (gene co-expression) does not automatically mean causation. This was confirmed to be true by successful simulation. Interestingly, when we separated the genes based on their chromosome location, they could be fitted with a third-degree polynomial, hinting toward the existence of linear correlation regulatory mechanisms. This suggests that gene expression falls into a conserved mechanism of expression. The simulation revealed that among different tumor samples,

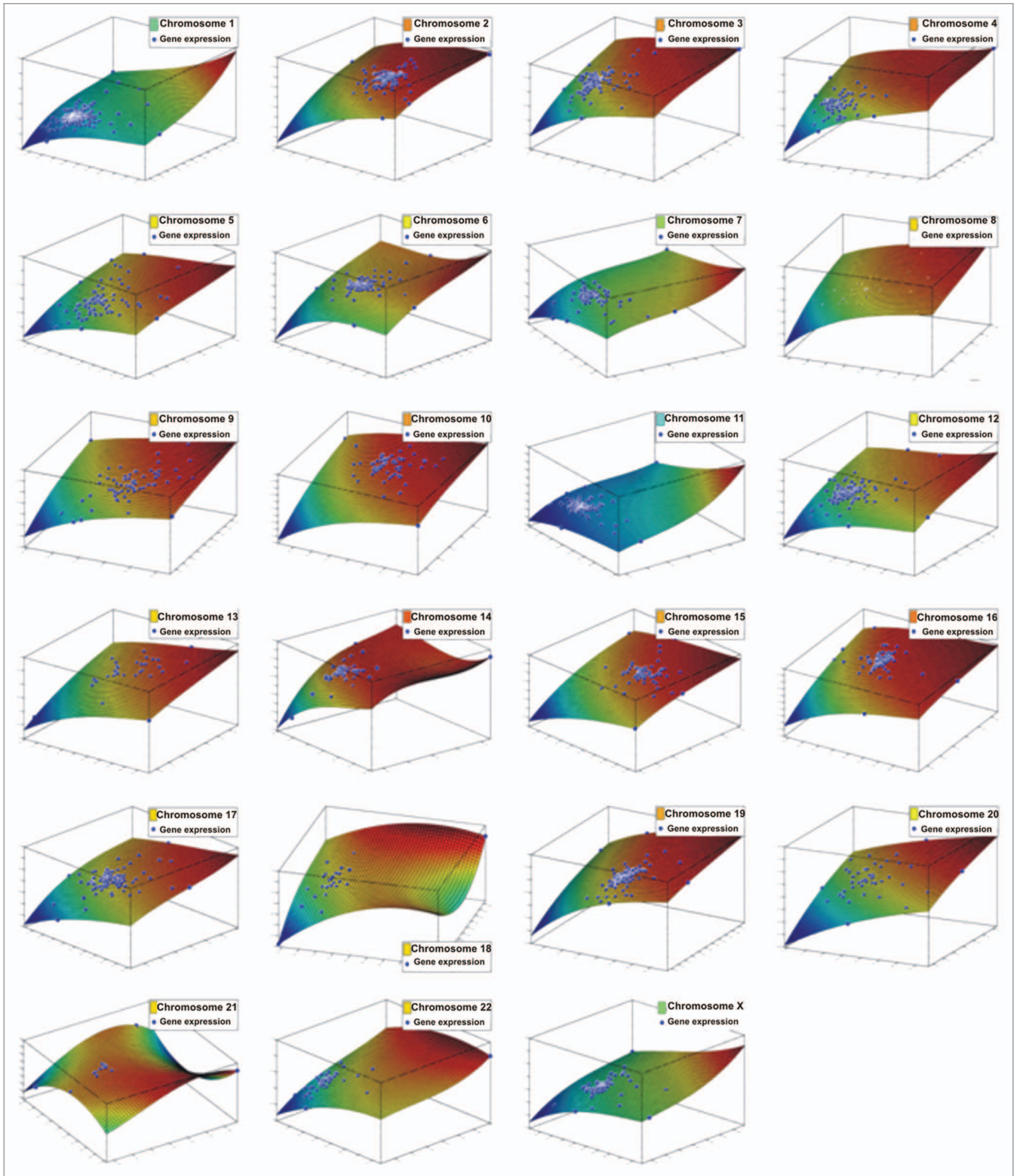
the genes on the same chromosomes were expressed in a similar manner.

Likewise, we constructed chromosome correlation maps for the T $\alpha$ -grade 2 tumors and found that chromosome-based gene expression manifested diverse correlation patterns. Thus, it was not a strong indication of correlation in this type of tumor. As in the case of T $\alpha$ -grade 1 tumors, expression data simulation also showed an excellent fit ( $0.7 < R^2 < 0.9$ ). These two cases suggest that correlation possibly also meant causation for these tumor types.

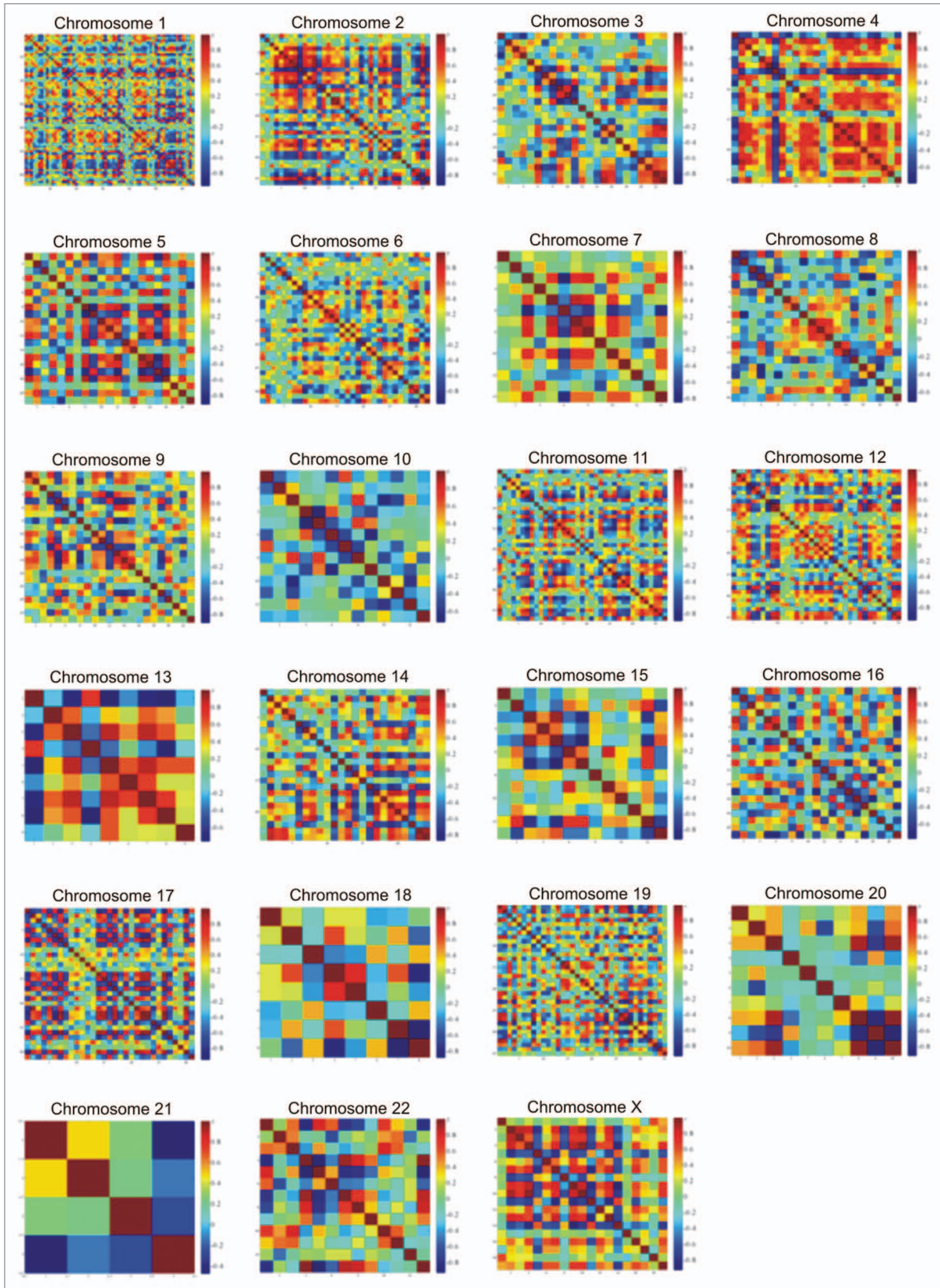
Next, we constructed chromosome correlation maps for the control samples alone in order to examine whether the correlations had to do only with relative ratios of tumors vs. the controls. Our analysis revealed chromosomal domains of gene co-expression for the control samples as well. Fitting attempts also showed that gene expression data could be simulated using polynomial functions, and again correlation hinted toward causation in normal tissue samples. To the best of our knowledge, this is the first time that microarray data are analyzed in such a way.

Despite the interesting type of mechanisms observed so far, especially in T $\alpha$ -grade 1 and T $\alpha$ -grade 2 tumors, we still wished to investigate whether such correlations appeared due to the small sample size that was used in the simulation procedures. Therefore, the next step was to perform the analysis in the complete sample size, in an attempt to gain insight into the whole picture of chromosome-based gene expression. For this purpose, we created the “gene cube.” The idea was that similarly regulated genes should manifest similar dynamics. Interestingly, it appeared that experimental data could be fitted with transformations, indicating that there are, at least in part, linear dynamics governing simplifications of the system described. New findings of molecular markers have been reported in the past in similar studies, using bioinformatics tools to identify gene expression signatures.<sup>25,26</sup> Thus the dynamics of chromosomal gene expression is of high significance in tumor biology. From our analysis, it was evident that gene expression has partly linear dynamics.

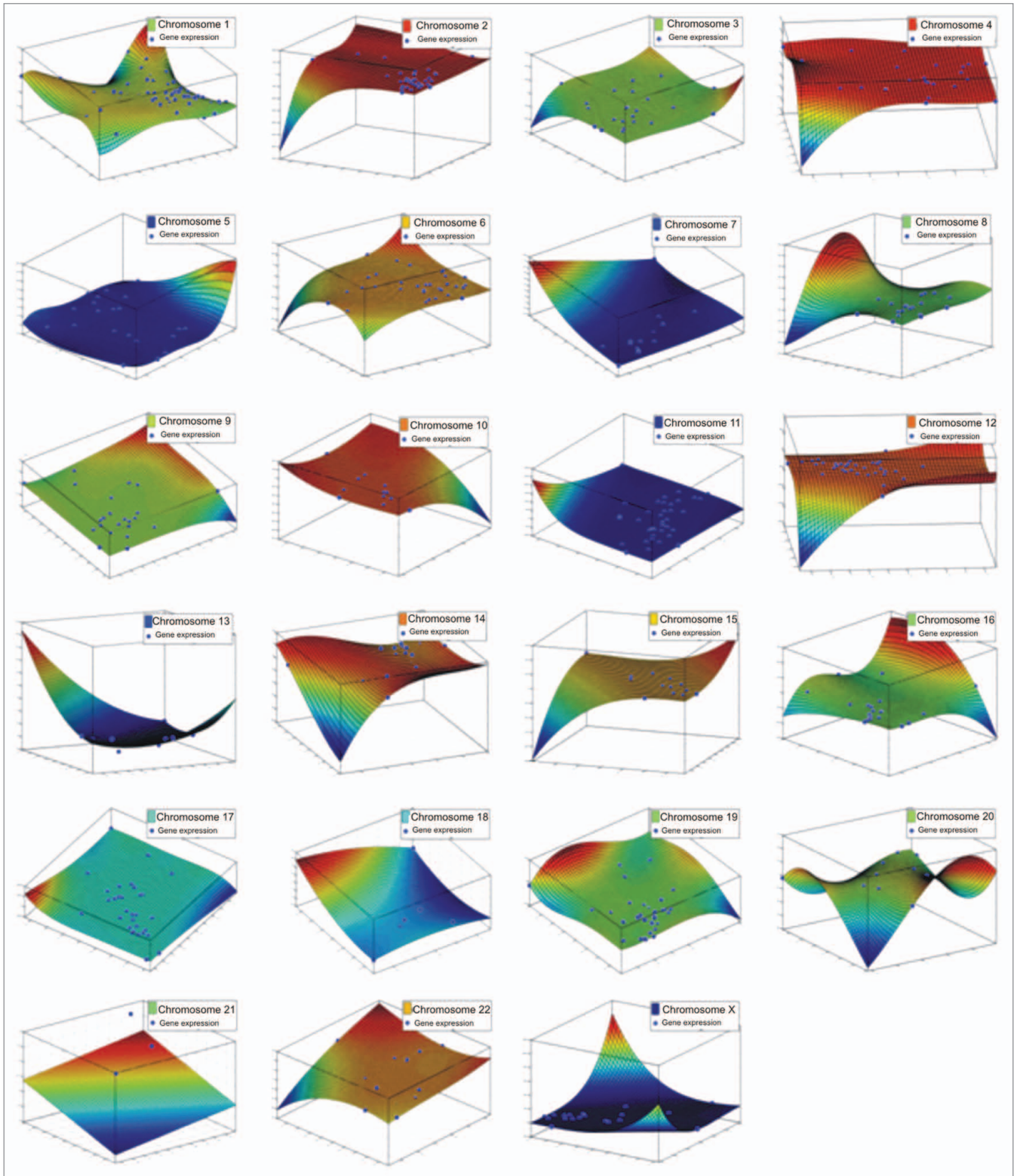
Genes on chromosomes 1 and 12 manifested functions related to embryonic development. The role of developmental genes in urothelial carcinomas was also previously reported.<sup>23,27</sup>



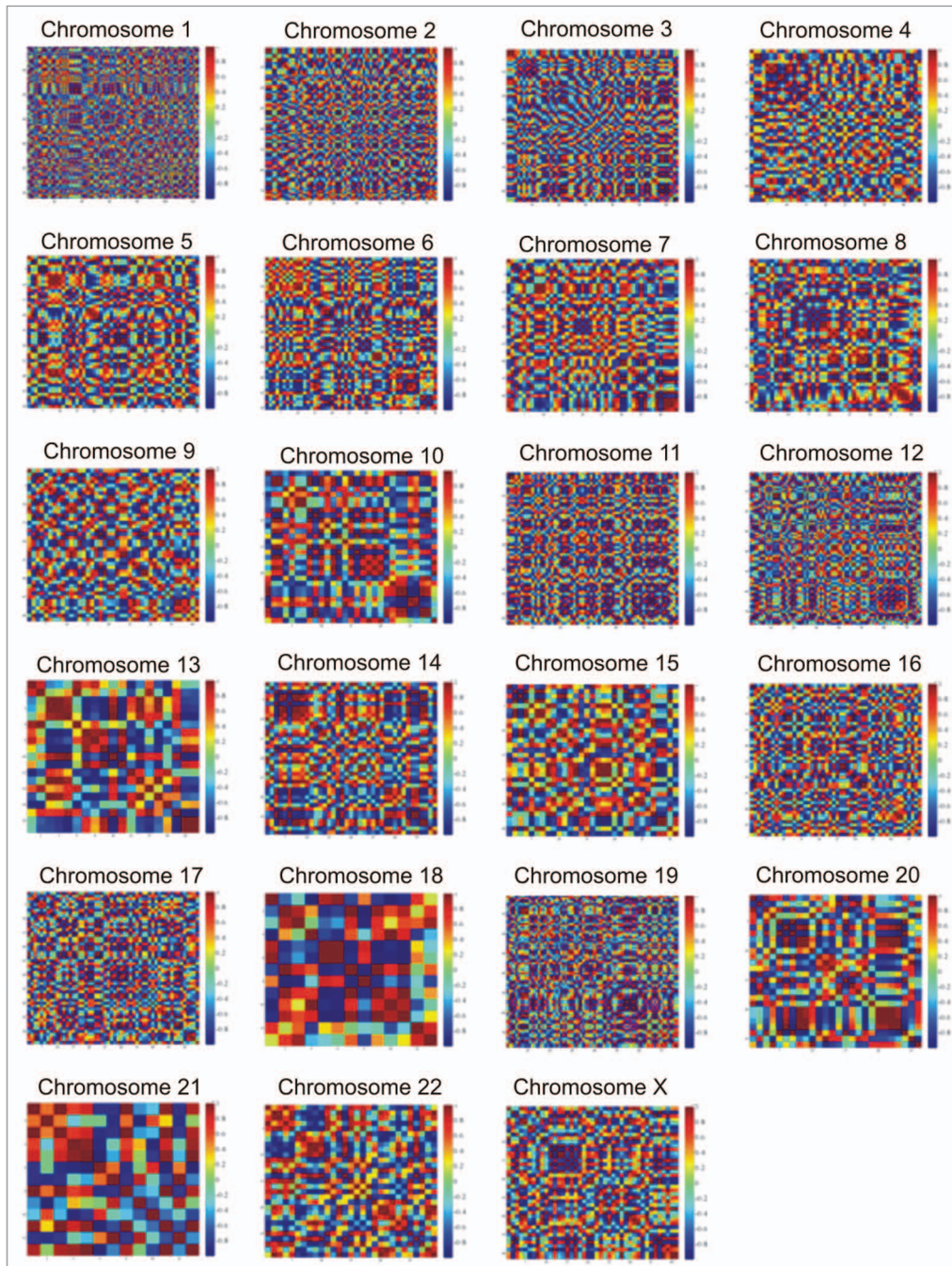
**Figure 4.** Simulations of the DE genes with respect to their chromosome location, among T $\alpha$ -grade 1 tumors and control samples. Each chromosome is presented separately. All genes could be simulated with a third-degree polynomial and  $R^2 > 0.99$ . Axes represent gene expression values of the  $\log_2$  ratio of the Ta-grade 1 tumors over control samples, where each axis represents one sample from the tumor subtype (Ta-grade 1 tumor group consisted of three samples).



**Figure 5.** Chromosome correlation maps for T $\alpha$ -grade 2 tumors allow visualization of co-expressed genes along all chromosomes.



**Figure 6.** Simulations of the DE genes with respect to their chromosome location, among T $\alpha$ -grade 2 tumors and control samples. Each chromosome is presented separately. All genes could be simulated with a third degree polynomial and  $R^2 > 0.99$ . Axes represent gene expression values of the  $\log_2$  ratio of the Ta-grade 2 tumors over control samples, where each axis represents one sample from the tumor subtype. Ta-grade 2 consisted of 12 samples in total. The figure includes representative fittings of the samples GSM2526\_Ta gr2, GSM2536\_Ta gr2 and GSM2507\_Ta gr2 for each chromosome, respectively.



**Figure 7.** Chromosome correlation maps for T1-grade 2 tumors allow visualization of co-expressed genes along all chromosomes. The X and Y axes represent the individual genes that were differentially expressed between control samples and T1-grade 2 tumors.

In particular, the genes related to this function were IL10 and VCAM1. VCAM1 has been reported to be upregulated in bladder carcinomas.<sup>28</sup> On the other hand, genes on chromosome 12 were LRP6, WNT1 and WNT10B. There are no reports linking *LRP6*, *WNT1* and *WNT10B* with bladder cancer. Genes on chromosome 10 were related to the WNT receptor signaling pathway (*BAMBI*, *BTRC*, *DKK1*, *FBXW4*, *FRAT1*, *FRAT2*, *FZD8*, *HHEX*, *LDB1*, *SFRP5*, *TCF7L2* and *WNT8B*). *BAMBI* has been linked with high-grade bladder carcinomas through epigenetic

silencing.<sup>29</sup> Also, *TCF7L2* has been reported to play a role in the WNT signaling pathway, where its inhibition seems to lead to G<sub>1</sub> arrest and growth inhibition of the bladder cancer. Genes on chromosome 4 and 9 participated in chemokine regulation. In particular, on chromosome 4, a large CXCL family is represented, including CXCL10, CXCL11, CXCL13, CXCL3, CXCL5, CXCL6, CXCL9 and IL8, PF4 and PPBP. The CXCL family of chemokines has been reported to participate in the microenvironment of bladder carcinoma.<sup>30</sup> IL8 has also been reported to

participate in bladder cancer.<sup>31</sup> *GHR*, *IL12B* and *PRLR* on chromosome 5 presented functions related to JAK2 kinase activity. So far, there are no reports connecting these genes with bladder tumors. Yet, it has been reported that the JAK/STAT participates in the inflammatory mechanisms of urinary bladder, relating it to the oncogenetic mechanisms of the disease.<sup>32</sup> Finally, on chromosome 22, MAPK1, MAPK11 and MAPK12 participate in the MAPK pathway. Interestingly, it has been reported that the inhibition of MAPK and NFκB signaling pathways inhibits growth and induces apoptosis of bladder tumor cells.<sup>33,34</sup>

Our data support the hypothesis that gene expression signatures can provide further information on gene regulatory mechanisms, based on chromosomal correlations of gene expression. The “Gene Cube” proved that gene expression has partly linear dynamics. Future research should focus on the investigation of these correlations in gene expression.

## Materials and Methods

**Strategy of the study.** In the first part, we performed microarray experimentation as previously described in detail<sup>7,35</sup> (Cohort A). The tumor samples were divided into three groups, as follows: T1-grade 2, T1-grade 3 and T2/T3-grade 3. All raw microarray data were MIAME compliant and uploaded on the GEO database (GSE27448).<sup>7</sup> The study was approved by the institutional review board of the University of Crete. For further data analysis, the Matlab<sup>®</sup> software was used.

In the second part, we extracted raw microarray expression data from 4 GEO data sets: (1) GSE89;<sup>13</sup> (2) GSE3167;<sup>14</sup> (3) GSE7476;<sup>36</sup> (4) GSE12630,<sup>37</sup> and incorporated them in our analysis, as previously described in detail.<sup>7</sup> All microarray data were background corrected and cross-normalized using a quantile algorithm.<sup>38-40</sup> In total, our pooled microarray analysis comprised 17 normal tissues and 129 bladder cancer samples (Cohort B).

All samples used from each data set with their respective GSM accession number, and detailed information regarding each sample’s tumor type has been previously described in detail.<sup>7</sup>

**Network analysis.** The co-deregulated genes among bladder cancer samples were investigated for network interrelation using the Ingenuity Pathways Analysis (IPA) software (www.ingenuity.com). The DE genes between cancer and normal tissue were used to generate a set of networks with a maximum network size of 35 genes/proteins. The median log<sub>2</sub> fold change value was used for analysis.

**Microarray data statistics.** First, we compared all the bladder cancer vs. all the normal tissue samples, entailing all bias by comparing them as unified groups. This analysis provided us with the co-DE genes. Second, we separated the 129 bladder cancer samples into 11 groups: (1) Tα-grade 1 tumors vs. controls; (2) Tα-grade 2 tumors vs. controls; (3) Tα-grade 3 tumors vs. controls; (4) T1-grade 2 tumors vs. controls; (5) T1-grade 3 tumors vs. controls; (6) T1-grade 2 and T1-grade 3 tumors vs. controls; (7) T1-grade 2, T1-grade 3 and T1-grade 4 tumors vs. controls; (8) group A<sup>15</sup> vs. controls; (9) group B<sup>15</sup> vs. controls; (10) group C<sup>15</sup> vs. controls; (11) metastatic tumors vs. controls. Each group was compared against all control samples. The false

discovery rate (FDR) was 8.6% for p < 0.05, calculated as previously described.<sup>41-43</sup>

**Cluster analysis.** K-means clustering with squared Euclidean as a metric distance was used to partition the gene expression profiles throughout the experimental setups.<sup>44</sup> The procedure was repeated 100 times, each with a new set of initial cluster centroid positions. The predictive power of the k-means algorithm was estimated using a figure of merit (FOM).<sup>45</sup> A FOM value vs. number of clusters was computed by removing each sample in turn from the data set, clustering genes based on the remaining data and calculating the fit of the withheld sample to the clustering pattern obtained from the other samples.

**Chromosome mapping and linear correlations.** Since consecutive genes are often similarly expressed, we mapped the genes on chromosome regions and identified their correlations through their chromosomal location.<sup>10</sup> The Gene Ontology Tree Machine, WebGestalt web-tool (Vanderbilt University, The Netherlands, <http://bioinfo.vanderbilt.edu/gotm/>)<sup>46</sup> and the Matlab<sup>®</sup> (The Mathworks Inc.) computing environments were used. Linear correlations were calculated using Pearson’s correlation coefficient. The value  $R^2 > 0.9$  was set as a threshold for statistical significance.

**Gene ontology (GO) enrichment.** GO analysis was applied to highlight the different functionalities among the experimental setups. For each chromosomal gene set formed, statistical analysis of GO term overrepresentation was performed against the common gene set, which was utilized as a reference set.<sup>47,48</sup> The chosen approach was the parent-child-union method.<sup>49</sup> Overrepresentation analysis (ORA) was performed with the publicly available Ontologizer 2.0 tool<sup>50</sup> using GO terms definitions and associations between genes and GO downloaded from the Gene Ontology consortium.<sup>51</sup> The threshold for statistical significance was set to < 0.01 using Bonferroni correction.

**Mathematical modeling and simulations.** Simulations were performed using Matlab<sup>®</sup>. For fitting purposes we used polynomial equations for 2D (Eqn. 1) and for 3D (Eqn. 2):

**Equation 1:**

$$f(x) = a_0 + a_1x^n + a_2x^{n-1} + \dots + a_nx$$

**Equation 2:**

$$f(x,y) = a_0 + a_1x^n + a_2x^{n-1}y + a_3x^{n-2}y^2 + a_4x^{n-3}y^3 + \dots + a_5x^3y^{n-3} + a_6x^2y^{n-2} + a_7xy^{n-1} + a_8y^n$$

Fourier series model as in Equation 3:

$$f(x) = a_0 + \sum_{i=1}^n (a_i \cos(mwx) + b_i \sin(mwx))$$

Sum of sin functions as in Equation 4:

$$f(x) = \sum_{i=1}^n a_i \sin(b_i x + c_i)$$

Fitting algorithm finds the coefficients of a polynomial  $f(x)$  of degree  $n$  that fits the data,  $f[x(i)]$  to  $y(i)$ , in a least squares sense. The result  $p$  is a row vector of length  $n+1$  containing the polynomial coefficients in descending powers as shown in Equations 1 and 2.

On the other hand the Fourier series is a sum of sine and cosine functions that describes a periodic signal (Eqn. 3). It is represented in either the trigonometric form or the exponential

**Table 3.** Selected GO terms of chromosomal-based gene expression

GO analysis		
ID	Name	p-value (Adj)
<b>chromosome 1</b>		
GO:0060669	embryonic placenta morphogenesis	0.0045
GO:0005839	proteasome core complex	0.0014
GO:0005031	tumor necrosis factor receptor activity	0.0067
<b>chromosome 2</b>		
GO:0009952	anterior/posterior pattern formation	0.0018
GO:0007350	blastoderm segmentation	0.0064
GO:0048468	cell development	0.0071
GO:0048562	embryonic organ morphogenesis	0.0025
GO:0048706	embryonic skeletal system development	0.0062
GO:0005152	interleukin-1 receptor antagonist activity	0.0052
GO:0004918	interleukin-8 receptor activity	0.0052
<b>chromosome 3</b>		
GO:0016493	C-C chemokine receptor activity	4.71E-05
GO:0071425	hemopoietic stem cell proliferation	0.0043
<b>chromosome 4</b>		
GO:0001525	Angiogenesis	0.0052
GO:0008009	chemokine activity	2.99E-07
GO:0008083	growth factor activity	0.0008
GO:0005134	interleukin-2 receptor binding	0.0059
GO:0051781	positive regulation of cell division	0.0001
GO:0045741	positive regulation of epidermal growth factor receptor activity	0.0008
GO:0050729	positive regulation of inflammatory response	0.0027
GO:0045410	positive regulation of interleukin-6 biosynthetic process	0.0003
GO:0045840	positive regulation of mitosis	0.0061
GO:0034105	positive regulation of tissue remodeling	0.0087
GO:0008202	steroid metabolic process	0.0065
<b>chromosome 5</b>		
GO:0042977	activation of JAK2 kinase activity	0.0010
GO:0035240	dopamine binding	0.0020
GO:0008083	growth factor activity	0.0048
GO:0070851	growth factor receptor binding	0.0056
GO:0005138	interleukin-6 receptor binding	0.0068
GO:0004923	leukemia inhibitory factor receptor activity	0.0023

The terms were considered significant if they obtained an adjusted p-value < 0.01.

form. The algorithm used in the present work used the trigonometric Fourier series form, where  $a_0$  models a constant (intercept) term in the data and is associated with the  $i = 0$  cosine term,  $w$  is the fundamental frequency of the signal,  $n$  is the number of terms (harmonics) in the series, and  $1 \leq n \leq 8$ .

**Table 3.** Selected GO terms of chromosomal-based gene expression (continued)

GO analysis		
ID	Name	p-value (Adj)
<b>chromosome 6</b>		
GO:0060333	interferon-gamma-mediated signaling pathway	1.78E-09
GO:0046415	urate metabolic process	0.0094
GO:0001570	vasculogenesis	0.0089
<b>chromosome 7</b>		
GO:0035425	autocrine signaling	0.0022
GO:0060571	morphogenesis of an epithelial fold	0.0072
GO:0045765	regulation of angiogenesis	0.0038
<b>chromosome 9</b>		
GO:0005125	cytokine activity	4.39E-05
<b>chromosome 10</b>		
GO:0005739	mitochondrion	0.0096
GO:0016055	Wnt receptor signaling pathway	0.0078
<b>chromosome 12</b>		
GO:0031076	embryonic camera-type eye development	0.0071
GO:0048048	embryonic eye morphogenesis	0.0051
<b>chromosome 16</b>		
GO:0006264	mitochondrial DNA replication	0.0068
GO:0005759	mitochondrial matrix	0.0060
GO:0010834	telomere maintenance via telomere shortening	0.0068
<b>chromosome 17</b>		
GO:0008009	chemokine activity	5.09E-05
<b>chromosome 21</b>		
GO:0004904	interferon receptor activity	0.0004
<b>chromosome 22</b>		
GO:0004707	MAP kinase activity	0.0007
<b>chromosome X</b>		
GO:0004896	cytokine receptor activity	0.0055

The terms were considered significant if they obtained an adjusted p-value < 0.01.

#### Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

#### Author's Contributions

G.I.L., A.Z. designed and supervised the study, performed the experimental procedures and data interpretation. A.Z. and G.I.L. prepared and submitted the manuscript. M.A., D.D., D.A. and S.V. read and approved the manuscript.

#### Supplemental Materials

Supplemental materials may be found here: [www.landesbioscience.com/journals/cc/article/24673](http://www.landesbioscience.com/journals/cc/article/24673)

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