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\*Manuscript

The application of artificial intelligence to microarray data: Identification of

a novel gene signature to identify bladder cancer progression

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1

### Abstract

Background: New methods to identify bladder cancer progression are required. Geneexpression microarrays can reveal insights into disease biology and identify novel biomarkers. However, these experiments produce large datasets that are hard to interpret.

Objective: To develop a novel method of microarray analysis combining two forms of artificial intelligence (AI): NeuroFuzzy Modeling (NFM) and Artificial Neural Networks (ANN). To validate this in a bladder cancer cohort.

Design, Setting, and Participants: We used AI and statistical analyses to identify progression-related genes in a microarray dataset (n=66 tumors, n=2,800 genes). The AI-selected genes were then investigated in a second cohort (n=262 tumors) using immunohistochemistry.

Measurements: We compared the accuracy of AI and statistical approaches to identify tumor progression.

Results and limitations: AI identified 11 progression-associated genes (OR=0.70 (95% CI=0.56-0.87) p=0.0004) and these were more discriminate than genes chosen using statistical analyses (OR=1.24 (95% CI=0.96-1.60) p=0.09). The expression of 6 AI-selected genes (LIG3, Fas, KRT18, ICAM1, DSG2 and BRCA2) was determined using commercial antibodies and successfully identified tumor progression (Concordance

Index=0.66, Logrank p=0.01). AI-selected genes were more discriminate than pathological criteria at determining progression (Cox multivariate analysis p=0.01). Limitations include the use of statistical correlation to identify 200 genes for AI analysis and that we did not compare regression identified genes with immunohistochemistry.

Conclusions: AI and statistical analyses use different techniques of inference to determine gene-phenotype associations and identify distinct prognostic gene signatures that are equally valid. We have identified a prognostic gene signature, whose members reflect a variety of carcinogenic pathways, which could identify progression in non-muscle invasive bladder cancers.

## Introduction

The care of patients with Urothelial Carcinoma of the bladder (UCC) could be significantly improved if their tumor behavior was accurately identified at diagnosis. Patients with non-progressive superficial disease could be spared endoscopic surveillance and BCG immunotherapy, whilst those at high progression risk could opt for early cystectomy. For invasive tumors the use of systemic chemotherapy could be rationalized to cases with highest progression risk. Tumor behavior can be hard to determine from histopathology alone. For example, the progression risk for non-muscle UCC varies between <1% and >50% [1, 2]. Furthermore, as stage and grade are often linked, when one is fixed (e.g. stage) the other performs poorly (e.g. grade) at identifying tumor progression. It is hoped that molecular knowledge will reveal an understanding of tumor biology that allows accurate phenotype identification.

As current biomarkers are insufficiently robust for clinical practice, microarrays have been used to identify new candidates [3] [4]. Microarray experiments reveal great insights into tumor biology but the cost and magnitude of these experiments prohibit large sample size analyses. Thus, microarray datasets have high dimensionality (large imbalance between gene number and sample size) that leads to analytical difficulties [5] [6] [7]. Successful analysis requires the identification of genes related to tumor-class and the removal of non-contributing variables. Poor analysis leads to data over-fitting and irreproducible results [5]. Traditional analytical techniques, such as hierarchical clustering, assume biological linearity and use statistical proximity to infer class-gene relationships (so called 'feature selection'). They perform poorly in datasets

contaminated with variable noise. Artificial intelligence (AI) is a machine learning approach without these prerequisites. Various AI techniques exist [8] and successful microarray analysis has been reported using artificial neural networks (ANN) [9] [10] and support vector machines (SVMs) [11, 12] in non-urothelial malignancies. However, the hidden working layer of an ANN prevents model understanding and hinders its acceptance by the scientific community [13], whilst SVMs still use proximity to infer class-gene associations and function poorly with respect to interpretability [14].

An alternative form of AI is the neurofuzzy model (NFM). This has a similar design to an ANN, but uses a transparent fuzzy logic internal structure [8]. This transparency allows model understanding, parameter interrogation and can facilitate the inclusion of priori qualitative knowledge. When used to identify tumor progression we have previously found that NFM is accurate, reproducible and appears superior to regression based classifications [15, 16]. We hypothesized that NFM could improve microarray analysis and identify prognostic gene panels that could accurately predict the behavior of UCC. To test this hypothesis we examined a previously reported non-muscle invasive UCC microarray dataset to find genes associated with progression to invasion. Genes associated with progression were then tested in a new larger UCC cohort using immunohistochemistry.

### Materials and methods

### Patients and Tumors

We studied two patient populations (Table 1). For microarray analysis we used 66 tumors from 34 patients, treated at the Ludwig Maximilian University, Germany (detailed in [17]). Progression to muscle invasion occurred in 10/34 patients (29%) and the median follow up was 43 months. For immunohistochemical analysis we studied 262 tumors from separate consecutive patients treated at the University of Regensburg, Germany. We created a tissue microarray (TMA) using paraffin embedded formalin fixed tissues with 2 cores per cases (1.2mm) [18]. Progression information was available for 182/262 (69.5%) patients and muscle invasion or new metastases occurred in 49 patients (26.9%). The median follow up was 89 months (range 2-154). No patients were in both UCC populations. Normal urothelium from patients with benign prostatic hyperplasia (n=20) and co-existing UCC (n=15) was also analyzed. Institutional review board approval was obtained from both institutions prior to study commencement.

## RNA Extraction and Gene Expression Microarray Analysis

The microarray (metg001A) contained 2,800 genes (6,117 probesets) annotated by the GoldenPath assembly. The microarray experiments and data processing are reported in detail elsewhere [17].

## Artificial Intelligence Feature Selection

To analyze the microarray data we used a 'Committee of models' approach that assimilated findings from each individual AI model (Figure 1), as we wanted to

determine gene-progression relationships that were not dependent upon one AI structure. We initially performed a dimension reduction using Pearson's coefficient to identify the 200 genes most associated with progression. These selected genes were then analyzed using iterative ANN and NFM models in two structures, which we termed 'Selectivity' and 'Averaging' (Figure 1). These structures enable simultaneous analysis of all genes, rather than a 'Leave-One-Out' approach. ANNs were produced within Statistica (Version 7, StatSoft Ltd, Bedford, UK). NFMs were produced within Matlab (Version 6.5 www.mathworks.com) and progression predictions performed using an in-house software suite [19, 20]. The data were divided into 90% for training (60% was learning and 30% for validation) and 10% for testing. Ensembling and cross validation were used to maximize data [21].

We ranked the 200 genes according to the size of model error induced by their alteration. Those with largest error were ranked highest, as alteration of their values produced the largest disturbance in the models accuracy. For each gene a 'Committee' ranking was produced from the average score of the individual AI models. A panel of progression related genes was produced from those with the highest ranking. This Committee panel was compared with the 'Original' gene panel selected using Pearson's linear regression coefficient and GeneCluster 2.0 software [17]. This Original panel included 11 members (FABP4, GSTM4, SERPINA1, HDAC1, C20ORF1, DNLC2A, PTK6 UBC, MGMT, ITGB3BP and PAIP2).

## *Immunohistochemistry*

To evaluate the *Committee* approach we analyzed the expression of its highest ranking members using immunohistochemistry in a new UCC cohort [17, 22]. Commercially manufactured antibodies were available for six members: LIG3 (clone 6G9; Abcam, Cambridge, UK; dilution 1:50), BRCA2 (Abcam, Cambridge, UK; dilution 1:10), TNFRSF6 (Abcam, Cambridge, UK; dilution 1:25), KRT18 (clone CK2; Chemicon, Billerica, MA, USA; dilution 1:50), DSG2 (clone 3G132; Abcam, Cambridge, UK; dilution 1:10), and ICAM1 (clone 23G12; Lab Vision, Fremont, CA, USA; dilution 1:10). For negative controls the primary antibody was omitted. Immunostained sections were scored independently for the percentage of positive tumor cells by uropathologists (PW, AH). The abnormal status for each protein was defined according to its cellular function, its contrast with normal urothelial expression and from previous reports. For ICAM1, a case was considered positive if > 30% of intra-tumoral blood vessels were stained. For LIG3, BRCA2, TNFRSF6, and DSG2 abnormal expression was defined as a loss or reduction of staining (0% or  $\leq$ 30% positively stained cells). For both, normal urothelium had expression in >50% of cells. Abnormal KRT18 expression was defined as increased immunostaining ( $\geq 80\%$  cells with positive staining) with respect to normal samples, which were negative in 90% of cases.

### Statistical Analysis

All analyses were two tailed and carried out using SPSS (version 14, SPSS Inc). Categorical variables were compared using the  $\chi 2$  test and continuous variables with a T test. Disease progression was defined when a non-muscle invasive tumor became invasive or a muscle invasive tumor developed metastases. Progression-specific survival

probability following tumor resection was analyzed using the Kaplan-Meier method and Log rank test. Patients without progression were censored when they were last reviewed or when they died of other causes. The concordance index was calculated as reported [23]. A P value of <0.05 was interpreted as statistically significant. Cox regression multivariate analysis was used to compare the prognostic value of the various gene panels with clinicopathological parameters.

### Results

#### Dimension reduction

We aimed to produce a prognostic gene panel of around 11 members to allow comparison with the *Original* panel chosen by statistical methods. Analysis of predictive ANN and NFM models with increments of 1 to 200 members revealed this was feasible (Figure 2). For NFM, the modeling error with 11 genes (RMS=0.135) was similar to that for more than 157 genes (both concordance index=1.0). For ANN the error did not change until more than 140 gene inputs were used (RMS = 0.37 for 11 genes), and was larger than the equivalent for NFM.

## Gene Ranking and Comparison of Feature Selection Panels

We ranked the 200 genes according to their average score from the various AI models (Table 2) and selected the 11 highest ranked genes to compare with the *Original* panel. Using gene expression, dichotomized around the mean, both panels were able to stratify tumor progression, although the *Committee* panel appeared more discriminate. For example, the findings of the *Committee* panel are typical (Figure 3a): whilst individual members are associated with tumor progression (e.g. LIG3 p=0.01, KRT18 p=0.04, Log rank values), the best prediction of progression occurs when the members are used in combination (≥3/11 abnormal genes p=0.007, ≥4/11 p=0.0004, ≥5/11 p=0.002, Log rank values). In multivariate analysis the *Committee* panel (OR=0.70 (95% CI 0.56-0.87), Logrank p=0.0004) was better at identifying progression than grade (OR=0.38 (95% CI 0.15-0.91, p=0.001) and stage (OR=0.65 (95% CI 0.1-4.31), p=0.03), and the *Original* 

panel (OR=1.24 (95% CI 0.96-1.60), p=0.09). No members were shared between the *Committee* and *Original* panels.

## Analysis of the Committee panel in a second tumor cohort

Six of the 11 members in the *Committee* panel (LIG3, BRCA2, TNFRSF6, KRT18, DSG2 and ICAM1, Figure 4a) have commercially manufactured antibodies with proven reproducible staining patterns in formalin fixed paraffin embedded tissue. Using these antibodies we performed immunohistochemistry on the 262 tumor TMA. When protein expression was analyzed with respect to tumor histology, various associations were seen. For example, LIG3 and ICAM1 were associated with tumor stage and grade ( $\chi^2$  p<0.05) (Table 3), when compared to tumors with normal expression. However, when expression of individual proteins with respect to tumor behavior was analyzed, few significant relationships were present. Only abnormal TNFRSF6 expression was significantly associated with tumor progression (Log rank p=0.003).

We then analyzed the 6 proteins together as a *Committee* panel using only superficial tumors (n=134). Each tumor was scored according to the number of proteins with abnormal staining and this was expressed as a percentage of the total number successfully immunostained for that sample. Only samples with ≥4 stained proteins were evaluated. When progression was analyzed with respect to this score, significantly worse outcomes were present in tumors with higher than lower scores (Figure 4b). As with its use in the first tumor cohort, the panel's discriminating ability was maximal at its mean content (Concordance index =0.66, Log rank p=0.02 for 40% and p=0.01 for 50%). In

multivariate analysis, the *Committee* panel was better at stratifying progression (Cox OR=1.2 (95% CI 1.1-1.3), p=0.014) than tumor stage (OR=1.44 (95% CI 0.82-2.53), p=0.2), grade (OR=0.93 (95% CI 0.53-1.66), p=0.8), the presence of CIS (OR=1.3 (95% CI 0.54-3.12), p=0.6), growth pattern (OR=0.74 (95% CI 0.26-2.12), p=0.6) and multifocality (OR=1.61 (95% CI 0.61-4.24), p=0.3).

### Discussion

Here we have used AI to examine the relationship between gene expression and progression. To evaluate this approach, rather than specific model designs, we used a Committee of models to merge gene rankings from individual models and structures. AI can identify complex relationships within non-linear data contaminated by variable noise and as such, can outperform statistical regression [8, 24]. AI modeling is a generic process and these methods could be applied to re-interrogate microarray datasets for prognostic and functional data.

Our approach reduced 200 genes to 11 with minimal deterioration in progression identification. The highest ranked genes appeared better at predicting tumor outcome than those selected using traditional analysis and pathological criteria. The fuzzy logic layer of our *Committee* NFM is shown in Figure 3b. This rule-base consists of parallel rules in which the fuzzy logic component can be visualized. In rule 1 (top line), high KRT18 in combination with low DSG2 and TNFRSF6 expression leads to rapid tumor progression (final box). This supports known carcinogenic functions of these genes as KRT18 is an oncogene and the others are tumor suppressors [25]. One can also see that the discriminatory effects in TP53BP2 are less apparent than for other genes (TP53BP2 was ranked 11th, Table 2).

The ability of AI to determine non-linear relationships is demonstrated in our results. Of the 11 genes that comprise the *Committee* panel, only TNFRSF6 was individually associated with tumor progression. However, the cumulative use of this panel allowed

accurate progression discrimination (Figure 4b). The members of the *Committee* panel represent various carcinogenic pathways. Their association with progression may be directly through carcinogenic roles or as bystanders associated with progression. Their diversity in roles suggests they may function as synergistic facilitators of progression. Apoptosis evasion is represented by reduced expression of Fas (TNFRSF6), TP53BP2 and ARHE. Fas is important for apoptosis induction and decreased expression is associated with advanced bladder cancer stage, grade and progression [26]. TP53BP2 (also 'Apoptosis stimulating protein of p53 2' (ASPP2)) plays a key role in apoptosis induction through the activation of p53. Reduced TP53BP2 expression abrogates the onset of apoptosis in cancer, but has not been reported in UCC. Tumor invasion is represented by reduced cellular adhesion (ICAM1 and DSG2) and cytoskeletal reorganization through increased KRT18 and reduced ARHE expression. DSG2 is a cellular adhesion molecule whose loss reduces adhesion, increases invasion and speeds tumor progression [27]. ICAM1 is also an intercellular adhesion molecule and is frequently epigenetically silenced in UCC (>70%) [28]. KRT18 is a cytokeratin known to be expressed in the umbrella layer of urothelium whose expression increases with urothelial carcinogenesis [25]. ARHE (also 'Rho family GTPase 3' (RND3)) is a Rho signal transduction member with roles in many cellular processes (cytoskeleton organization, membrane trafficking, cell growth and apoptosis) [29], whose loss is reported in prostate cancer. Deranged DNA repair is represented by BRCA2 and LIG3 [30]. Whilst neither is directly linked with bladder carcinogenesis, it is possible that loss of both is required for carcinogenic alteration. BRCA2-deficient cells have reduced DNA ligation capacity which can be reversed by LIG3 administration [30].

## Conclusion

AI can analyze microarray datasets in a complementary manner to statistical analyses. Both methods use different techniques of inference to determine gene-phenotype associations and thus identify distinct prognostic gene signatures that are equally valid. We have identified a new prognostic gene signature in UCC, whose members reflect a variety of carcinogenic pathways. This signature requires validation in new tumor cohorts to assess its ability to identify progression in non-muscle invasive bladder cancers.

## Conflict of interest

We declare no conflicts of interest with this work.

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## Figure Legends

Fig. 1. The Work flow for this report. Pearson coefficient was used to reduce the 2,800 genes to the 200 most associated with progression. These genes were modelled by separate ANNs and NFMs. For each model 200 iterations were run. Each iteration studied a single gene and consisted of training/validation/testing of the model. The model's error was the score that determined the significance of the gene being tested in that iteration. In the *Selectivity* approach, we changed all 200 gene values to their mean and then individually maximized (largest value seen) and minimized single genes. Following model testing the analysed gene was returned to average before starting the next iteration. This approach hoped to find genes whose extreme presence caused most disruption to the model. In the Averaging approach, all 200 genes were left unchanged whilst single individual genes were averaged for their model iteration. This model aimed to find those genes whose loss of profile resulted in most disruption to the model. Gene rankings from these models were then averaged to generate the Committee of models ranking. The highest ranking members were then compared with the Original panel identified by Wild et al [17] by predicting progression in the same UCC cohort. Six members had commercial antibodies and their expression was tested in a new cohort of UCC.

Fig. 2. Performance of the AI models during dimension reduction. (a). The model error for 1-200 genes is shown (RMS value). In general, NFM has a lower error than ANN. A panel with 11 genes has a similar error to that with 158 genes (NFM) or 140 genes

(ANN). (b) Correct progression classifications (percentage) with NFM or ANN using models with  $n^{1-200}$  genes.

Fig. 3. The *Committee* panel for Superficial UCC progression prediction. (a). Tumour progression stratified by pathological grade, the *Original* and *Committee* panels. (b). NFM rule base for the *Committee* panel. Probe values are coloured according to value around mean (reduced = red, increased = blue and mean = black).

Fig. 4. Tumor progression using immunohistochemistry for members of the *Committee* panel. (a) Abnormal expression of panel members in UCC (Insert boxes show expression in normal urothelium: strong membranous TNFRSF6 staining; KRT18 confined to umbrella cells; strong nuclear LIG3 and BRCA2 staining; weak membranous DSG2 staining; very weak ICAM1 staining of urothelium and strong staining within the endothelium of capillaries). (b). Tumor progression stratified by grade and the *Committee* panel in the second superficial UCC population (n=134). A bad signature is defined as ≥50% proteins with abnormal expression.

Table 1. The two UCC cohorts studied in this report

		i. Gene arra	ay tumors	ii. TMA tumors				
		n	%	n	%			
Gender	Male	29	85.3%	194	73.8%			
	Female	5	14.7%	68	25.9%			
Stage TNM 1998	Normal	8	100.0%					
	pTis	3	4.5%					
	рТа	46	69.7%	149	56.7%			
	pT1	10	15.2%	49	18.6%			
	pT2	7	10.6%	59	22.4%			
	pT3			2	0.8%			
	pT4			3	1.1%			
Stage TNM 2004	PUNLMP	1	1.5%	22	8.4%			
	pTis	3	4.5%					
	рТа	45	68.2%	127	48.3%			
	pT1	10	15.2%	49	18.6%			
	pT2	7	10.6%	59	22.4%			
	pT3			2	0.8%			
	pT4			3	1.1%			
Grade	Grade 1	27	40.9%	83	31.6%			
	Grade 2	24	36.4%	69	26.2%			
	Grade 3	15	22.7%	110	41.8%			
Growth pattern	Papillary	55	83.3%	210	79.8%			
	Solid	11	16.7%	51	19.4%			
	Unknown			1	0.4%			
Multiplicity	Unifocal	29	43.9%	54	20.5%			
	Multifocal	37	56.1%	208	79.1%			
Carcinoma in situ	No pTis	62	93.9%	227	86.3%			
	pTis	4	6.1%	35	13.3%			
Tumor	Metastasis			2	0.8%			
	Primary UCC	25	37.9%	255	97.0%			
	Recurrent UCC	41	62.1%	5	1.9%			
	Progression rate	10/34* 29.4%		36/134***	26.9%			
Median (range) t	21 (0-60)	months	23 (1-154) months					
	Overall survival	unknown		167/198	<i>*</i>			
Median (range) over	43 (0-109)	) months	90 (24-154)	(24-154) months				
	Total UCC	66	100.0%	262	100.0%			

<sup>\*</sup> in the 34 individual patients

<sup>\*\*</sup> in non-progressing patients

<sup>\*\*\*</sup> in 134 primary superficial tumors with available follow up information

Table 2. The *Committee* gene panel selected according to ranking frequency from AI models

	Symbol	Gene name	Function
1	PP	Pyrophosphatase (inorganic)	Phosphate metabolism / metabolism
2	TNFRSF6	TNF receptor family member 6 (CD95)	Apoptosis / immune response / signal transduction
3	LIG3	DNA Ligase III	Cytokinesis / DNA replication & repair / meiosis
4	BRCA2	Breast cancer type 2 susceptibility gene	Cell cycle control / double-strand break repair / DNA replication / chromatin architecture / apoptosis
5	ICAM1	Intercellular adhesion molecule 1 (CD54)	Cell-cell adhesion
6	ARHE	Ras homolog gene family, member E	Cell adhesion / signal transduction / cytoskeleton organization
7	NACA	nascent-polypeptide-associated complex $\alpha$	Protein biosynthesis / nascent polypeptide association
8	DSG2	Desmoglein 2	Cell adhesion / homophilic cell adhesion
9	KRT18	Keratin 18	Embryogenesis and morphogenesis
10	FLJ14146	Uncharacterized protein C1orf115	Unknown function
11	<i>TP53BP2</i>	p53 binding protein 2 (ASPP2)	Cell cycle/ apoptosis regulation / signal transduction

Genes shown in bold were analysed by immunohistochemistry

Table 3. Immunohistochemical analysis of 263 bladder tumors for L/G3, BRCA2, TNFRSF6, KRT18, DSG2 and /CAM1

	TNFRSF6			LIG3			ICAM1		DSG2			BRCA2			KRT18			
	Abnorm	nal / To	otal χ2	Abnorm	al / To	otal χ2	Abnorm	al / To	otal χ2	Abnorm	al / To	tal χ2	Abnorn	nal / To	otal χ2	Abnorn	nal / To	tal χ2
Grade																		
1	16/69	28%		42/66	64%		46/61	75%		35/67	52%		23/69	33%		29/61	48%	
2	15/65	23%		44/66	67%		41/62	66%		37/65	57%		22/66	33%		30/60	50%	
3	13/93	14%	0.093	33/95	35%	0.0001	40/93	43%	0.0001	63/93	68%	0.119	40/95	42%	0.398	46/88	52%	0.850
Stage																		
PUNLMP	7/19	37%		10/18	56%		11/16	69%		12/18	67%		5/18	28%		7/16	44%	
рТа	28/112	25%		70/110	64%		72/103	70%		55/109	51%		37/113	33%		51/100	51%	
pT1	6/42	14%		17/43	40%		26/44	59%		31/43	72%		18/42	43%		21/41	51%	
pT2	6/49	12%		21/51	41%		18/48	38%		34/50	68%		23/52			24/48	50%	
pT3	0/2	0%		0/2	0%		0/2	0%		1/2	50%		0/2	0%		2/2	100%	
pT4	0/3	0%	0.119	1/3	33%	0.018	0/3	0%	0.001	2/3	66%	0.128	2/3	67%	0.344	0/2	0%	0.506
CIS																		
Absent	42/200	21%		110/198	69%		115/189	61%		117/169	60%		75/201	37%		90/182	50%	
Present	5/27	19%	0.765	9/29	31%	0.014	12/27	44%	0.105	18/29	62%	0.807	10/29	35%	0.768	15/27	56%	0.554
Growth																		
Papillary	42/187	23%		99/185	53%		115/177	63%		103/184	56%		69/187	37%		88/170	52%	
Solid	4/39	10%	0.085	20/41	49%	0.583	14/38	37%	0.003	31/40	78%	0.012	15/42	36%	0.886	17/38	45%	0.433
Progression free	survival	*																
Progression	12/34	35%	0.003	21/33	64%	0.566	17/31	55%	0.155	23/33	70%	0.081	13/33	40%	0.802	17/30	57%	0.458
No	10/84	12%		48/83	58%		54/78	69%		42/81	52%		31/84	37%		35/72	49%	

<sup>\*</sup> Only superficial tumors were analyzed for this outcome

NOTE. For each variable the numerator is the number of abnormally immunostained tumours and the denominator is the number successfully analyzed for that protein

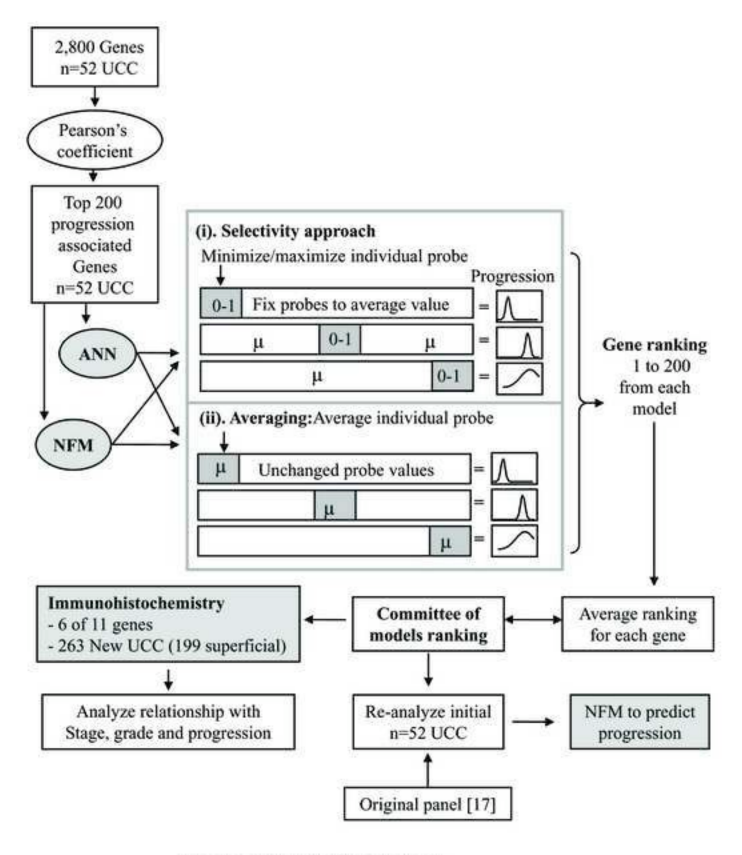


Figure 1. Work flow for this report.

Figure 2 Click here to download high resolution image

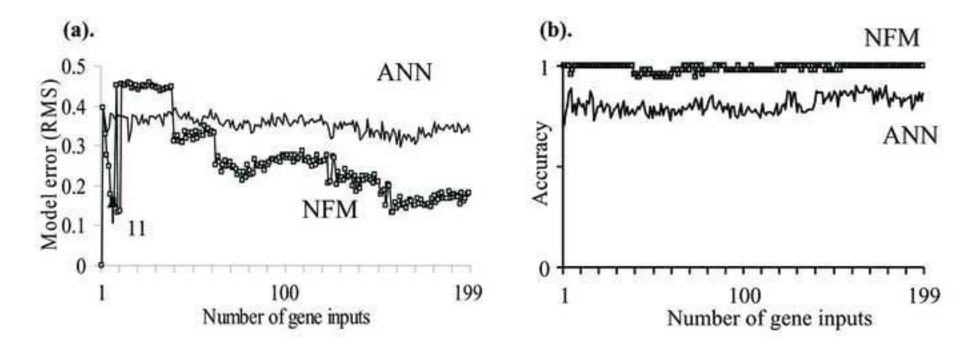


Figure 2. Performance of the AI models during dimension reduction.

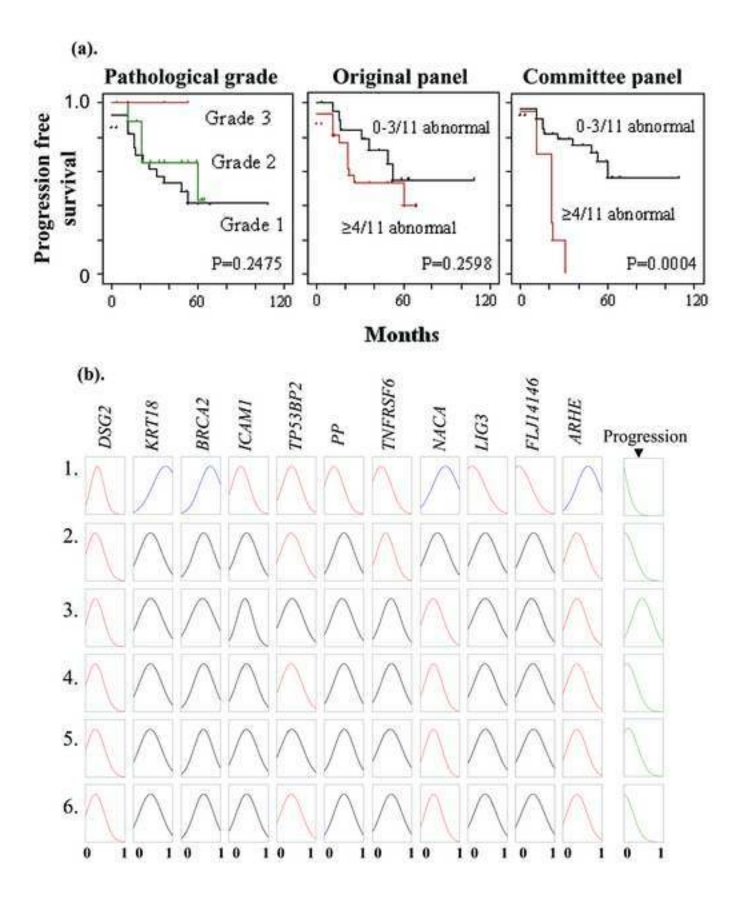
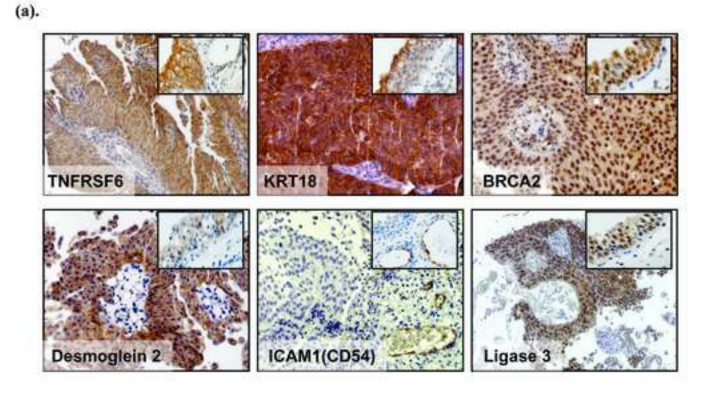


Figure 3. The Committee panel for Superficial UCC progression prediction.

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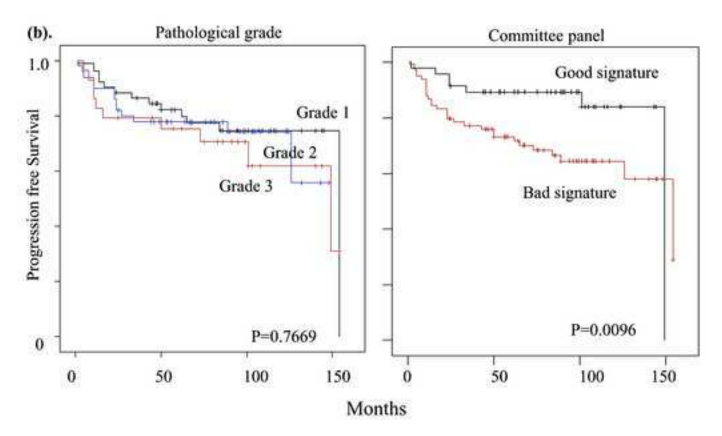


Figure 4. Tumor progression using immunohistochemistry for members of the Committee panel.