

Building an Outcome Predictor Model for Diffuse Large B-Cell Lymphoma

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Diffuse large B-cell lymphoma (DLBCL) patients are treated using relatively homogeneous protocols, irrespective of their biological and clinical variability. Here we have developed a protein-expression-based outcome predictor for DLBCL. Using tissue microarrays (TMAs), we have analyzed the expression of 52 selected molecules in a series of 152 DLBCLs. The study yielded relevant information concerning key biological aspects of this tumor, such as cell-cycle control and apoptosis. A biological predictor was built with a training group of 103 patients, and was validated with a blind set of 49 patients. The predictive model with 8 markers can identify the probability of failure for a given patient with 78% accuracy. After stratifying patients according to the predicted response under the logistic model, 92.3% patients below the 25 percentile were accurately predicted by this biological score as “failure-free” while 96.2% of those above the 75 percentile were correctly predicted as belonging to the “fatal or refractory disease” group. Combining this biological score and the International Prognostic Index (IPI) improves the capacity for predicting failure and survival. This predictor was then validated in the independent group. The protein-expression-based score complements the information obtained from the use of the IPI, allowing patients to be assigned to different risk categories. (*Am J Pathol* 2004, 164:613–622)

Diffuse large B-cell lymphoma (DLBCL) is the most frequent type of lymphoma, with a 5-year survival probability

of around 50%. Although a significant proportion of DLBCL patients can be cured with current combination chemotherapy regimes, at present there is no clinical or biological score available that can accurately distinguish patients who can be cured with standard therapy and those who require new treatment approaches.¹

Outcome with DLBCL, as in other types of cancer, is the result of interactions between the genetic abnormalities in the tumor and the clinical status of the patients. Information concerning the molecular abnormalities present in DLBCL, derived from genome-wide expression analysis, allows us to identify multiple markers that suggest the existence of a vast number of underlying genetic events in all of the major cell pathways involved in control of proliferation, apoptosis, signal transduction, DNA repair, and other processes.^{2–4} Nevertheless, until recently, outcome-predictor systems have been based on single genetic abnormalities, or the integration of clinical data into predictive models, such as the International Prognostic Index (IPI).⁵

Tissue microarrays (TMAs) are a powerful and reproducible technique for demonstrating the biological variability inherent in cancer and, when applied to lymphoma samples, are capable of identifying multiple alterations in the regulation of critical genes and pathways.^{6,7}

In the present study we have investigated the expression of a large number (52) of markers in a DLBCL series using TMAs. The results yield information concerning the variety of molecular markers that predict clinical response. These can be integrated into a single predictive model that identifies the probability of failure with 78% accuracy. This biological score can be used to complement the information obtained by the use of the IPI, allowing patients to be stratified into different risk categories.

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Materials and Methods

DLBCL Samples

235 cases of DLBCL were collected. These were diagnosed between 1990 and 1999, the stages being evaluated according to standard protocols. Patients were treated with regimes including polychemotherapy (mainly adriamycin-based) with or without adjuvant radiotherapy and/or surgery. Diagnostic paraffin blocks were selected on the basis of the availability of suitable formalin-fixed paraffin-embedded tissue, containing enough remaining tissue as for a minimum of 60 sections. Histological confirmation of DLBCL was achieved in all cases by central review using standard tissue sections. Histological criteria used for diagnoses and classification of cases were those described in the World Health Organization classification.⁸ Paraffin-embedded blocks from reactive lymphoid tissue, cell lines and different B- and T-cell lymphoma samples, used for control purposes, were obtained from the tissue archives of the CNIO Tumor Bank.

Tissue Microarray Design

We used a Tissue Arrayer device (Beecher Instruments, Sun Prairie, WI) to construct three different TMA blocks, containing 502 cylinders in total, according to conventional protocols.⁷ All cases were histologically reviewed and the most tumor-rich areas were marked in the paraffin blocks. Two selected 0.6-mm-diameter cylinders from two different areas were included in each case, along with 16 separate controls to ensure the quality, reproducibility and homogenous staining of the slides. Selected controls include reactive lymph nodes and tonsils, and paraffin-embedded cell lines.

Immunohistochemical staining was performed and evaluated for the 50 different antibodies using standard procedures.⁷ The selected markers correspond to sets of key proteins involved in cell cycle, apoptosis (extrinsic and intrinsic pathways), and B-cell differentiation, additionally including a large majority of the markers previously identified as survival predictors in DLBCL.

Staining of TMA sections was evaluated by three different pathologists (A.S., J.F.G., F.C.), using uniform criteria. To guarantee the reproducibility of this method, we decided to employ straightforward and clear-cut criteria. After initial analysis, the pattern of staining for each Ab was recorded as positive *versus* negative, or high *versus* low level of expression, taking into account the expression in reactive and tumoral cells and specific cut-offs for each marker. Specific details of the threshold used in each case are given in Table 1. As a general criterion, these thresholds were preferentially selected on the basis of their reproducibility and, when possible, their ability to correlate with previous findings using these markers and/or specific biological events.

As cytoplasmic STAT1, STAT3, and NF κ B expression can generally be found in normal lymphoid cells and lymphomas, we have considered as positive cases only those showing distinct nuclear expression in the

tumoral cells, thereby indicating the activated form of these proteins.⁹

Discrepancies between the two cylinders included for each case were resolved through a reviewed joint analysis of both cores. The same procedure was applied to discrepancies among pathologists.

The reactivity of most of the antibodies used here has been validated in previous studies.⁷

In situ detection of apoptosis and EBER *in situ* hybridization (ISH) were performed using standard procedures,⁷ using the appropriate controls. Apoptosis was detected using the ApopTag Peroxidase In Situ Apoptosis Detection Kit (Intergen Co., Oxford, UK). Epstein-Barr virus (EBV) was detected by ISH with fluorescein-conjugated Epstein-Barr Virus (EBER) PNA probe (DAKO, Glostrup, Denmark). EBV-positive cases were considered to be those showing EBER nuclear expression in a majority of the tumoral cells.

Validation of the Technique

The reproducibility of the results obtained was confirmed by comparing them with those from whole sections from 42 randomly selected cases that had been stained using the same procedures for a selection of markers including CD20, bcl-2, and bcl-6.

Statistical Study

The Pearson χ^2 statistic and the Spearman correlation coefficient were used as appropriate to analyze relationships between the 52 markers studied.

Survival analyses were performed on all patients for whom follow-up information was available for a minimum of 24 months (approximately 70% of the overall series) and who had complete expression analysis data. HIV-positive patients⁹ were excluded from the outcome analysis. The final number of patients included in the survival analysis was 152, all of them treated with curative intention.

Failure was defined as the absence of complete remission, progression, or death attributable to the tumor. The series was divided into a training group of 103 cases for the purpose of building the predictor, and a second, smaller group of 49 cases, to validate the model.

Overall Survival (OS) and Failure-Free Survival (FFS) curves were plotted using the Kaplan-Meier method. Statistical significance of associations between individual variables and OS or FFS was determined using the log-rank test.

Cox's univariate proportional hazard analysis was also performed independently for each variable. Results were validated by multiple testing and the random permutation test.

For multivariate analysis, the series was divided into a training group of 103 cases for the purpose of building the predictor, and a second, smaller group of 49 cases, to validate the model.

A logistic regression model was used to predict failure. Only variables identified in the univariate analysis asso-

Table 1. Antibodies Used in the Analyses, Indicating Source, Dilution, Threshold and Pattern of Reactivity Used and Positive Controls

Protein	Clone	Source	Dilution	Reactivity	Threshold	Internal control
Bcl-2	124	DAKO	1:25	High/low	>50% positive cells	Small lymphocytes
Bax	POLYCLONAL	Santa Cruz	1:1000	Positive/negative	>10% positive cells	Benign B lymphocytes
Bcl-XL	2H12	ZYMED	1:10	High/low	>10% positive cells	TMA controls
Mcl1	POLYCLONAL	DAKO	1:100	High/low	>50% positive cells	Proliferating cells
Survivin	POLYCLONAL	RD Systems	1:1500	High/low	>10% positive cells	TMA controls
p65/RelA	F-6 (p65)	Santa Cruz	1:2000	Positive/negative	Nuclear expression	TMA controls
Caspase 3	C92-605	PharMingen	1:25	Positive/negative	>10% positive cells	TMA controls
Bcl-10	331.3	Santa Cruz	1:1000	Positive/negative	>10% positive cells	Reactive lymphocytes
CD95	GM30	Novocastra	1:50	Positive/negative	>10% positive cells	Reactive lymphocytes
Oct-1	12F11	Santa Cruz	1:10	Positive/negative	>10% positive cells	Reactive lymphocytes
Oct-2	POLYCLONAL	Santa Cruz	1:500	Positive/negative	>10% positive cells	Reactive lymphocytes
Bob-1	POLYCLONAL	Santa Cruz	1:3000	Positive/negative	>10% positive cells	Reactive lymphocytes
PU1	G148-74	PharMingen	1:50	Positive/negative	>10% positive cells	Benign B-lymphocytes
Pax-5	POLYCLONAL	Santa Cruz	1:200	Positive/negative	>10% positive cells	CG (germinal center) B cells
MUM-1	POLYCLONAL	Santa Cruz	1:200	High/low	>80% positive cells	Plasma cells
STAT3	F-2	Santa Cruz	1:500	Positive/negative	Nuclear expression	Reactive lymphocytes, macrophages
Bcl-6	PG-B6p	DAKO	1:10	Positive/negative	>10% positive cells	CG (germinal center) B cells and B-cell lymphomas
CD38	VS38	DAKO	1:25	High/low	>80% positive cells	Plasma cells
CD138	MI15	DAKO	1:50	High/low	>80% positive cells	Plasma cells
CD5	4C7	Novocastra	1:50	Positive/negative	>10% positive cells	Reactive lymphocytes
CD10	56C6	Novocastra	1:10	Positive/negative	>10% positive cells	CG (germinal center) B cells
CD20	L-26	DAKO	1:100	Positive/negative	Any positive tumoral cell	Reactive lymphocytes
CD30	15B3	Novocastra	1:100	Positive/negative	>10% positive cells	TMA controls
EMA	E29	DAKO	1:50	Positive/negative	>10% positive cells	TMA controls
CD27	137B4	Novocastra	1:150	Positive/negative	>10% positive cells	Reactive lymphocytes
Cyclin A	6E6	Novocastra	1:100	Positive/negative	>10% positive cells	Proliferating cells (G2/M)
Cyclin B1	7A9	Novocastra	1:25	Positive/negative	>50% positive cells	Proliferating cells (G2/M)
Cyclin D1	DCS-6	DAKO	1:100	Positive/negative	Any positive tumoral cell	Macrophages and endothelial cells
Cyclin D3	DCS-22	Novocastra	1:10	Positive/negative	>50% positive cells	Proliferating cells
Cyclin E	13A3	Novocastra	1:10	High/low	>80% positive cells	TMA controls, proliferating cells
CDK1	1	Transduction	1:1500	Positive/negative	>80% positive cells	TMA controls, proliferating cells
CDK2	8D4	NeoMarkers	1:500	Positive/negative	>50% positive cells	TMA controls, proliferating cells
CDK6	K6.83	Chemicon	1:10	Positive/negative	>80% positive cells	TMA controls
P21	EA10	Oncogene	1:50	Positive/negative	>10% positive cells	Scattered GC cells
P16	POLYCLONAL	Santa Cruz	1:50	High/low	>10% positive cells	Normal cells
P27	57	Transduction	1:1000	High/low	>10% positive cells	Resting lymphoid cells
Ki67	MIB1	DAKO	1:100	High/low	>50% positive cells	Proliferating cells
SKP2	1G12E9	ZYMED	1:10	Positive/negative	>80% positive cells	Proliferating cells
P53	DO-7	Novocastra	1:50	High/low	>80% positive cells	Scattered GC cells
Hdm2	IF2 (Mdm2)	Oncogene	1:10	High/low	>10% positive cells	Macrophages, endothelial cells
Rb	G3-245	BD PharMingen	1:250	High/low	>80% positive cells	Proliferating cells
Rb-P (Phospho-Rb)	sc-7986-R	Santa Cruz	1:250	High/low	>80% positive cells	Proliferating cells
PTEN	28H6	Novocastra	1:50	Positive/negative	Any positive tumoral cell	Normal cells
DP-1	1DP06	NeoMarkers	1:50	Positive/negative	>80% positive cells	Proliferating cells
PKCβ	28	Serotec	1:500	Positive/negative	>10% positive cells	Plasma cells, endothelial cells
TOPO	Ki-S1	DAKO	1:400	High/low	>50% positive cells	Proliferating cells
GST	353-10	DAKO	1:150	High/low	>50% positive cells	Proliferating cells
c-kit	POLYCLONAL	DAKO	1:25	High/low	>10% positive cells	Stromal cells
ALK	ALK1	DAKO	1:50	High/low	>10% positive cells	TMA internal controls
CD3	F7.2.38	DAKO	1:25	Positive/negative	Any positive tumoral cell	Reactive lymphocytes

DAKO, Glostrup, Denmark; Santa Cruz Biotechnology, Santa Cruz, CA; BD PharMingen, San Diego, CA; Novocastra, Newcastle, UK; Transduction Laboratories, Lexington, KY; Neomarkers, Fremont, CA; Chemicon, Temecula, CA; Oncogene Research Products, Darmstadt, Germany; Serotec, Oxford, UK.

ciated with FFS with values of $P < 0.2$ and in which at least 5 cases were considered positive or negative were included. Highly variable components in the model were excluded, since they could have introduced uncertainty in predictions. For comparative purposes, multivariate models using step-up (forward) variable selection and other heuristic procedures were also fitted. The final model estimates values of the odds ratio (OR), 95% confidence interval (CI) and P for each variable. General

applicability of the model was tested by leave-one-out cross-validation. The stability of the model was evaluated by influence statistics (DfBeta). Different predictor models were found, when using the leave-one-out cross-validation, but these showed only small variations in the weight of each marker, or selection of markers. Accuracy was also tested by the Receiver Operating Characteristic (ROC) curve, which allows the discriminating ability of the model to be estimated.

To demonstrate the predictive capacity of the model, patients were ranked according to this score and then divided into four equal groups, or quartiles. To validate the model overall, the specific weight or coefficient assigned to each gene, as determined in the preliminary group, was applied to calculate the outcome-predictor score in the validation group. Once the model had been validated, a final logistic regression model was fitted to the entire data, allowing adjustment of the coefficients. Statistical analyses were performed using the SPSS program and the tools at <http://bioinfo.cnio.es/> for random permutation tests.

Results

The percentage of informative individual cores was 90.4%. As each TMA included 2 different core cylinders from each marker, the final percentage of missing expression data values was 12% (Table 2).

To check the reliability and accuracy of TMA for this measure of protein expression, TMA and quantitative whole tissue stainings were compared in a subset of 42 cases. Concordances of 100%, 91.1%, and 90% were obtained for CD20, bcl-2 and bcl-6, respectively, thus coinciding with the results of other NHL analysis studies.^{10,11}

Results of the overall DLBCL series are summarized in Table 2. Figure 1 shows the expression of the markers found to predict failure after the multivariate analysis.

Correlation between Markers

The Pearson test revealed a large number of significant associations between the different markers analyzed. Full details of the correlation between markers are given in Supplementary Appendix 2 at http://bioinfo.cnio.es/data/DLBCL_TMA.

The most striking findings were as follows:

- Higher levels of expression of specific cyclins and CDKs were observed in varying numbers of this series: 11.7% (25 of 214) in the case of cyclin E, 52.9% (197 of 202) for cyclin A, 22.3% (40 of 179) in the case of CDK1 and 76.3% (151 of 198) for CDK2. There was a close relationship between proliferation and apoptosis, including the positive association observed between proliferation and apoptotic indices, and between the apoptotic index and different CDKs and cyclins.

- EBV presence was accompanied by changes in the expression of numerous proteins, including an increase in CDK1, cyclin B1, SKP2, p21, CD30, and a loss of BOB1, pax-5, and bcl-6.

- SKP2 expression, observed in 12.1% (26 of 214) of cases, was significantly associated with changes in numerous apoptosis and cell-cycle regulators, including a strongly positive correlation with CDK1, Rb, cyclin A, B1, D3, survivin, and a negative association with Bax and bcl-2. A significant relationship was also observed between SKP2 expression and the increased expression of Rb-P, CDK6, MDM2, p53, bcl-6, CD10, c-kit, EBER, NF-κB, caspase 3 active, MCL1, MIB1, and TUNEL.

Table 2. Expression of 51 of the 52 Markers Analyzed in the Entire DLBCL Series, Indicating Number of Positive/Total Analyzed Cases

Protein	Positive cases	Percentage
Apoptosis		
MBcl-2	122/224	54.46
Bax	194/215	90.23
Bcl-XL	73/188	38.83
Mcl1	107/186	57.53
Survivin	70/217	32.26
p65/RelA	116/225	51.56
Caspase 3 active	17/194	8.77
Bcl-10	75/188	39.89
CD95	69/169	40.83
TUNEL	141/191	73.82
Transcription factors		
Oct-1	186/187	99.46
Oct-2	189/192	98.44
Bob-1	176/180	97.78
PU1	11/194	5.67
Pax-5	209/215	97.21
MUM-1	113/206	54.85
STAT3	23/224	10.27
B-cell differentiation		
Bcl-6	168/207	81.16
CD38	73/204	35.78
CD138	15/219	6.85
CD5	53/188	28.19
CD10	51/182	28.02
CD20	224/231	96.97
CD30	41/206	19.90
EMA	8/214	3.74
CD27	30/209	14.35
Cell cycle		
Cyclin A	107/202	52.97
Cyclin B1	37/221	16.74
Cyclin D1	0/235	0
Cyclin D3	50/229	21.83
Cyclin E	25/214	11.68
CDK1	40/179	22.35
CDK2	151/198	76.26
CDK6	111/174	63.79
P21	20/226	8.85
P16	166/212	78.30
P27	78/216	36.11
MIB1	131/210	63.38
SKP2	26/214	12.15
p53	37/222	16.66
Hdm2	120/221	54.30
Rb	112/215	52.09
Rb-P	57/189	30.16
Other		
PTEN	211/211	100
DP-1	114/155	73.55
PKCβ	55/186	29.57
TOPO	186/207	89.85
GST	150/209	71.77
c-kit	60/213	28.17
ALK	1/213	0.47
EBER	20/221	9.05

CD3 was negative in all cases.

- An unexpected finding was the association of c-kit expression with various cell cycle markers (increased p27, SKP2, CDK1, cyclin E, and Rb-P), apoptosis (loss of Bax and increase in bcl-XL, bcl-10, survivin), high PKCβ,

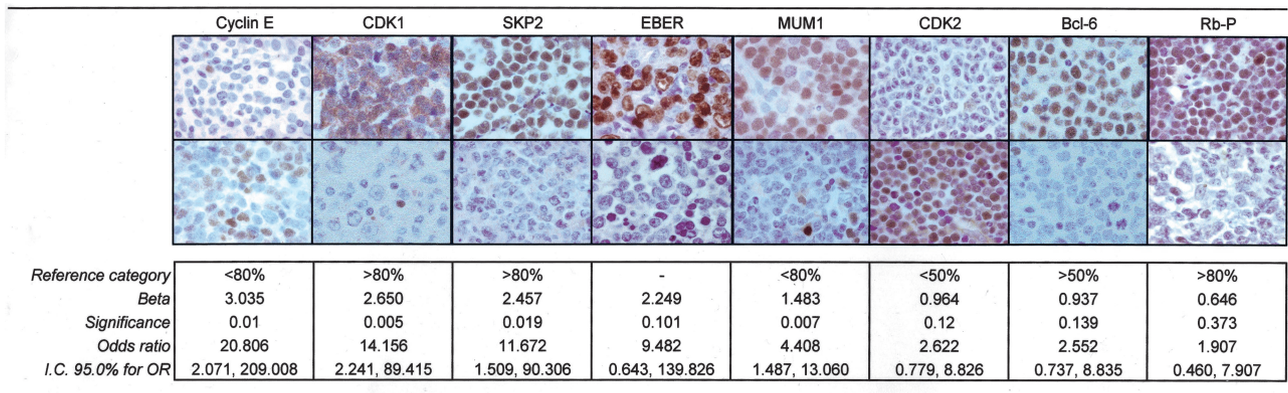


Figure 1. Characteristics and variables included in the biological model for failure prediction in DLBCL. Representative immunohisto chemistry and *in situ* hybridization results for the eight markers selected in the multivariate analysis. A positive and a negative tumor are shown for each marker. β represents the weight of each variable estimated from the multivariate analysis.

and B-cell differentiation (elevated CD27, CD38, CD5, CD10).

• Finally, bcl-6 expression, detected in 81% (169 of 207) of cases, was associated with profound changes in molecules regulating cell cycle (high SKP2, CDK6, MDM2, Rb, Rb-P, and loss of p21), apoptosis (increase in bcl-x_L and NF-kB), and B-cell differentiation (increase in Pax5, CD10, and Bob1 expression). Notably, it was found to be inversely correlated with EBV and epithelial membrane antigen (EMA). Another interesting finding was the existence of a group (47% of cases) that simultaneously expressed bcl-6 and MUM1, two proteins that normal lymphoid B cells do not express at the same time.

Correlation between Protein-RNA Expression and Outcome in DLBCL

To detect any possible selection bias, the 152 included patients (Table 3) were compared with those who had been excluded due to insufficient follow-up. Comparison of age, gender, clinical stage and IPI revealed no significant differences.

All 52 individual variables were analyzed using Kaplan-Meier plots and Cox proportional hazard models to determine whether the expression was significantly associated with changes in FFS (Table 4). Ten variables (cyclin E, CDK1, SKP2, bcl-6, p21, Oct-2, BOB1, EMA,

Bax, bcl-2) were significantly correlated with FFS ($P < 0.05$) and nine showed a non-significant trend ($P < 0.2$). All of the significantly FFS-correlated variables, except Rb-P, were also associated with OS probability ($P < 0.05$) (data not shown). Furthermore, EBER, which showed a non-significant trend in the FFS analysis, was found to be associated with OS ($P < 0.05$) (data not shown). The result of the Cox proportional hazard analysis was then validated using multiple testing and random permutation tests ($n = 1000$).

Predicting Failure in DLBCL

Logistic regression analysis was used to find a DLBCL outcome predictor, making it possible to recognize which patients could be cured by the application of chemotherapeutic regimes. The group of 103 cases was used to build the predictor. Only variables identified in the univariate analysis associated with FFS with values of $P < 0.2$, and in which at least five cases were considered positive or negative, were included (19 variables, excluding EMA, Oct-2, BOB1). The final logistic regression model included the following markers: cyclin E, CDK1, SKP2, EBER, MUM1, CDK2, bcl-6, and Rb-P (Figure 1).

The predictor is a biological score, the probability of "failure" for one patient, which is calculated as

$$P = \frac{1}{(1+e^{-z})}$$

where

$$z = -7.0865 + 3.0352 \times \text{cyclinE} + 2.6502 \times \text{CDK1} + 2.4572 \times \text{SKP2} + 2.2494 \times \text{EBER} + 1.4833 \times \text{MUM1} + 0.9639 \times \text{CDK2} + 0.9367 \times \text{bcl-6} + 0.6458 \times \text{Rb-P}$$

and where coefficients from the logistic model are used as weights for the corresponding markers.

The percentage of correct classification for this model, using the training series, was 78.64% (81.13% for predicting FFS and 76% for patients with treatment failure).

Table 3. Clinical Characteristics of the 152 DLBCL Patients Included in the Outcome Analysis

Age (mean, range)		58.4 (5–96)
Gender	Female	47.6%
	Male	52.4%
IPI	0–1	41.8%
	2	27.1%
	3	17.1%
	4–5	14.1%
Follow-up (median)		60.9
Overall survival	5-year cumulative survival	59.8%
Failure	Cured versus fatal/refractory disease	50.6%/49.4%
Failure-free survival	5-year cumulative survival	50.4%

Table 4. Univariate Analysis for OS and FFS in the Current Series (*n*: 152 Patients) and Logistic Regression Model for Failure Prediction in the Training Set of Patients (*n*: 103)

Protein	Reference category	Univariate analysis for FFS (Cox regression)				Multivariate analysis for failure protein, RNA-expression-based model and model integrating IPI; (logistic regression)		
		<i>P</i>	RR	95% CI		Beta in PEB model	Beta in PEB + IPI model	Difference between models
				Lower	Higher			
IPI	IPI (0–2)	0.000	3.257	2.121	5.001		3.260	
cyclin E	<80%	0.000	3.293	1.839	5.894	3.035	2.477	0.184
CDK1	>80%	0.029	2.281	1.090	4.771	2.650	2.975	–0.123
SKP2	>80%	0.019	3.999	1.261	12.683	2.457	2.329	0.052
EBER	–	0.086	1.898	0.913	3.945	2.249	2.569	–0.142
MUM1	<80%	0.071	0.065	0.409	1.037	1.483	1.758	–0.185
CDK2	<50%	0.114	0.623	0.347	1.120	0.964	0.739	0.233
Bcl-6	>50%	0.040	1.747	1.027	2.972	0.937	0.655	0.301
Rb-P	>80%	0.117	1.648	0.882	3.078	0.646	1.037	–0.391
p21	–	0.001	3.042	1.601	5.780			
cyclin B1	>50%	0.094	1.869	0.899	3.883			
cyclin A	>50%	>0.2						
MDM2	–	0.097	1.446	0.936	2.234			
Rb	>80%	0.192	1.342	0.863	2.088			
CD38	<80%	0.188	1.369	0.858	2.184			
CD138	<80%	0.090	1.882	0.906	3.907			
Oct_2*	+	0.015	4.270	1.325	13.757			
BOB1*	+	0.003	8.836	2.074	37.657			
EMA*	–	0.040	2.891	1.052	7.948			
CD95	–	>0.2						
Bax	–	0.037	3.428	1.080	10.879			
Bcl-2	<50%	0.015	1.740	1.114	2.719			

*, <5 values in one category; –, no data available.

PEB, protein-expression-based; RR, relative risk.

Specific weight (beta) of each variable for predicting failure in the protein and RNA-expression-based model, compared with values for model integrating the IPI. Differences were calculated as (beta1-beta2)/beta1.

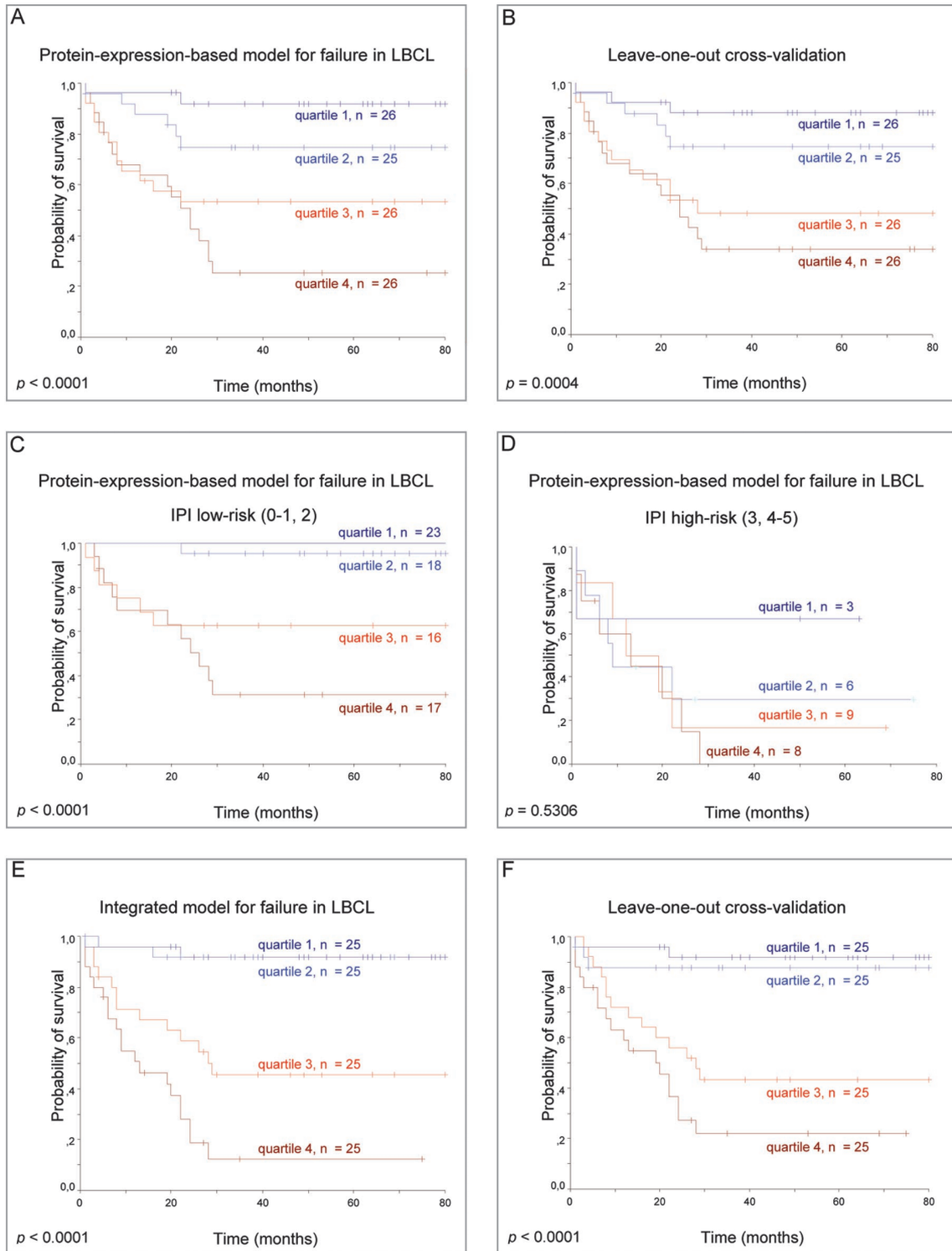
In a second step, patients were ranked according to their protein-expression-based score (0 to 1) and divided into four different quartiles, according to their specific risk. Stratifying patients according to these quartiles, 92.3% of patients beneath the 25 percentile were accurately predicted as “failure-free” by the score, and 96.2% of the patients above the 75 percentile were correctly predicted as belonging to the group of “fatal or refractory disease”. Between the 25 and 75 percentiles the accuracy of prediction fell below 90% for both categories (64% in the second quartile and 53.8% in the third quartile). Thus, when assigning each patient a specific risk, the capacity for predicting the upper and lower quartile is much higher than for patients with intermediate quartiles.

Validating the Biological Score for Failure in DLBCL

A Kaplan-Meier survival analysis, classifying patients according to the quartile of assigned probability, confirmed that the patients predicted to be cured had significantly improved long-term survival compared with those predicted to have fatal/refractory disease (5-year OS: 91.97% below the 25 percentile, vs. 25.45% above the 75 percentile; *P* < 0.0001) (Figure 2A).

The prediction accuracy of the score was then assessed using a leave-one-out cross-validation testing method, withholding one case and using the remaining set of tumors to train the model, predicting the “failure” probability of the withheld case. The process was repeated until all 103 samples had been predicted in turn. The results confirmed, with minor differences, the FFS and OS predictive capacity of the biological score (Figure 2B). Different predictor models were found, when using the leave-one-out cross-validation, showing only small variations in the weight of each marker, or selection of markers.

Although the majority of the patients of this series received anthracycline-based chemotherapy, 12 of 103 (11.6%) patients were treated with different drugs. To examine whether the biological model was independent of the treatment regimes used, treatment was included as a new variable. The specific weight of each variable in the model remained similar (3.064 × cyclin E + 2.499 × CDK1 + 2.364 × SKP2 + 2.264 × EBER + 1.391 × MUM1 + 1.088 × CDK2 + 0.898 × bcl-6 + 0.828 × Rb-P). Moreover, the correct classification percentage in this new model with the variable “treatment” decreased imperceptibly (77.2% for the overall prediction). Correct prediction percentage in the different quartiles was 92% (quartile 1 for



1. Quartile of protein-expression-based score (A) and leave-one-out cross-validation (B).
2. Quartile of protein-expression-based score for each IPI category (C and D).
3. Quartile of protein-expression-based and IPI score (E) and leave-one-out cross-validation (F).

Figure 2. Protein-expression-based model for failure prediction in DLBCL. Kaplan-Meier estimation of OS according to the assigned probability for each model in the training set of patients. Quartiles of protein-expression-based score (A) and leave-one-out cross-validation (B). Quartiles of protein-expression-based score for each IPI category (C) and leave-one-out cross-validation (D). Quartiles of protein-expression-based and IPI score (E) and leave-one-out cross-validation (F).

failure-free) vs. 96.2% (quartile 4 for failure). These percentages are very similar to those obtained previously.

Integration of Protein-Expression-Based Score and IPI

This biological score yielded a 13.616-fold odds ratio (OR) [95% CI (5.288, 35.063), $P < 0.0001$] for failure of treatment (percentile 50). IPI (low risk *versus* high risk), the standard clinical score for predicting the outcome in DLBCL,⁵ in this series yielded a 10.151-fold OR [95% CI (3.159, 32.616), $P < 0.0001$] for failure. A multivariate analysis including both the IPI and the protein-expression-based score showed that the significance of the biological score for failure [percentile 50; OR = 18.983; 95% CI (5.988, 60.180); $P < 0.0001$] seemed to be superior to and independent of the IPI [OR = 15.359; 95% CI (3.672, 64.244); $P < 0.0001$].

To determine whether the information contained in the protein and RNA-expression-based model was the same as or additional to the variables included in the IPI, patients were classified into low-risk (IPI: 0–2) and high-risk groups (IPI: 3–5), and then the protein-expression-based score quartiles were used in both groups. Low-risk IPI patients were accurately stratified by the protein-expression-based score into groups with a failure probability of 95.24% (quartile 4), 81.89% (quartiles 3 and 2) and 31.59% (quartile 1), $P < 0.00001$. High-risk IPI patients were also discriminated into two main groups using the protein-expression-based score, although the difference was not significant. These results suggest that an integrated use of the IPI and the protein-expression-based score could improve the predictive capacity of the model (Figure 2, C and D).

The joint predictive capacity of the protein-expression-based score and IPI was analyzed in a multivariate model. The specific weight of each component of the biological score in this new model remained quite similar (Table 4), confirming that the biological and clinical scores contain at least partially independent information. The predictive capacity of the model incorporating the IPI and the variables integrated in this biological score was slightly higher than that based purely on the protein and RNA-expression-based model, with 83% overall correct classification of failure (92% for quartile 1 and 96% for quartile 4).

This was correlated with a better discrimination of patients with different outcomes. Thus, patients allocated above the 50 percentile of the integrated score had 91.73% 5-year OS *versus* 29.71% for patients predicted for “failure” (Figure 2E).

Blind Test for Validation of the Predictor

The leave-one-out cross-validation confirmed the high predictive capacity of this integrated model, with a probability of failure in each respective quartile of 12%, 24%, 68%, and 88%, reflected in the overall survival probability (Figure 2F). The discriminating ability of this model was better than that of the protein and RNA-expressed-based

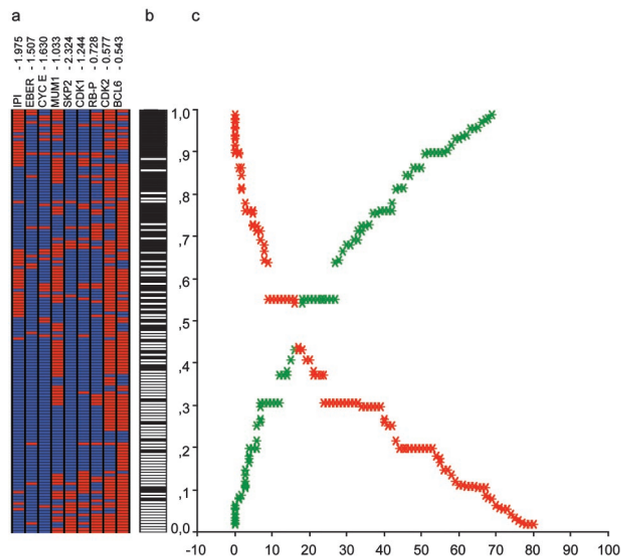


Figure 3. Final biological and clinical predictor model. **a:** Tree-view representation of the eight markers and IPI. Each column represents a marker, while each row corresponds to a patient, ordered according to the assigned failure probability. Specific weight of each marker is included at the top of each column. **b:** Real status of each patient (failure, black *vs.* maintained complete response, white). **c:** Graphic representation of the relation between the assigned probability and the real status. The graphic represents the accuracy of the predictor model. If the probability assigned to each patient (*y* axis) is less than 0.5, the model classifies the case into the group of maintained response. If the probability is greater than 0.5, the system classifies the case as a failure. The curves represent the number of patients erroneously classified as failure (in red), and those cases erroneously predicted to maintain a complete response (in green). Eventually, a threshold for each curve of cumulative error could be chosen to select a group of patients with a high probability of failure or of maintained complete remission.

model [ROC curve area: 0.901; $P < 0.0001$, 95% CI (0.840, 0.961)].

As this evaluation was based on the same training set of patients from which the predictive model was derived, we decided to estimate the accuracy of the classifier with an additional cohort of 49 patients who had not previously been included. In this independent series, the failure prediction and the outcome were evaluated by the model integrating the 8 markers and IPI, using the threshold from the training set of patients. The immunostaining and evaluation of these tumors were performed independently of the previous cases. The predictive capacities of the validation and preliminary group were comparable with respect to the assigned score for each patient by the model (76.9% and 83.3% of correct classification into quartiles 1 and 4, $P < 0.001$). Furthermore, values for 5-year OS were closely related with the assigned failure probability for each patient (5-year OS: 100%, 81.48%, 75%, and 25% for each quartile of the score; $P < 0.0001$).

Once the model had been validated, a final model with the 8 biological markers and IPI was fitted to the entire data (training + validation series). Finally, the biological-IPI score allowed assignment of a case-specific probability of failure, as can be observed in Figure 3.

Discussion

DLBCL seems to be the result of deregulation of multiple genes involved in the control of cell cycle, apoptosis, cell

growth, DNA repair, ubiquitin degradation, and other processes. Particularly striking is the existence of multiple concurrent abnormalities in the genes and pathways in the control of cell cycle and apoptosis. Subtle alterations in this exquisitely regulated balance between cell proliferation and apoptosis seem to contribute critically to DLBCL pathogenesis.

Some of the observed changes affect the large majority of cases analyzed here, such as the expression of bcl-6. The hypothetical relevance of bcl-6 in DLBCL pathogenesis is underlined by the increasing number of bcl-6 targets that are being described in B cells, and for its capacity to contribute to oncogenesis by rendering cells unresponsive to antiproliferative signals from the p19(ARF)-p53 pathway, as demonstrated by Shvarts et al.¹² In this respect, it is noteworthy that in this series bcl-6 expression appears to be associated with down-regulation of p21 and overexpression of MDM2. The potential role of bcl-6 as a promoter of cell-cycle progression beyond the G1/S restriction point is suggested by the existence of an additional significant relationship with increased phosphorylated Rb. Our data also confirm the prognostic significance of bcl-6 expression in DLBCL, as previously pointed out, when taking into account bcl-6 mRNA expression levels.¹³

According to the results of this study, Skp2 expression, which increased in one-fifth of the cases analyzed, is associated with many changes in apoptosis and cell-cycle regulators. Protein degradation throughout the ubiquitin pathway thus seems to be indicated as a potential contributory factor in the deregulation of proliferation and apoptosis in DLBCL.^{14,15} In addition to the confirmed role of Skp2 for inducing the degradation of p27 and Cdk2-unbound cyclin E, an accelerated degradation of unknown additional substrates is likely to play a role in oncogenic events mediated by Skp2.¹⁵

Cyclin E overexpression is highlighted by the uni- and multivariate analyses as a clinically highly relevant adverse prognostic marker, thus confirming previous observations in specific lymphoma types^{16,17} and other tumors.¹⁸ A possible explanation for these findings is provided by the recent demonstration that overexpression of cyclin E leads to increased chromosome instability and impaired S-phase progression.¹⁹

In general, the results of the univariate analysis confirm those previously published concerning single markers, such as the case for bcl-2 or others.^{20,21} Nevertheless, some of the significant markers in the univariate analysis, can prove not significant in the multivariate analysis.

Results of this study, not based on previous hypotheses of DLBCL subclassification, are difficult to match with the three DLBCL subgroups defined by Rosenwald et al⁴: germinal-center B-cell-like, activated B-cell-like, and type 3 diffuse large B-cell lymphoma. Instead, it seems that the tumors accumulate alterations in critical pathways stochastically, leading to the increased proliferation and loss of apoptosis observed here. The existence of a large group of double bcl-6+ MUM1+ cases demonstrates that the mutual exclusion of these markers, as observed in reactive germinal centers, is not preserved in

DLBCLs.²² Tumoral cells probably take advantage of the simultaneous expression of both proteins.

The technique used here is based on large-scale analysis of protein expression, detected by immunohistochemistry. The use of tissue microarrays is limited by the relatively small number of markers chosen (52 in this case), although it has the advantage of using protein profiling, which probably reflects more closely the characteristics of the tumoral cells than does RNA detection.

The integration of these markers into a single model allows the assignment of a specific probability of failure to each patient, according to the biological and clinical characteristics of each case. This information could eventually be used for individualized treatments, in which patients are stratified into therapeutic groups. A clinical application of this and other studies should, nevertheless, first fulfill the necessity of demonstrating the reproducibility of immunohistochemistry techniques among different groups, which would be facilitated by the application of automated systems for scoring immunohistochemical expression.

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