A "GENIUS" Approach for Breast Cancer Prognostication

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- Head: Prof. Christos Sotiriou.
- 8 researchers (1 prof, 5 postDocs, 2 PhD students), 5 technicians.
- Research topics : Genomic analyses, clinical studies and translational research.
- Website :

http://www.bordet.be/en/services/medical/array/practical.htm.

- National scientific collaborations : ULB, Erasme, ULg, Gembloux
- International scientific collaborations : Genome Institute of Singapore, John Radcliffe Hospital, Karolinska Institute and Hospital, MD Anderson Cancer Center, Netherlands Cancer Institute, Swiss Institute of Bioinformatics, NCI/NIH, Gustave-Roussy Institute, IDDI.

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- Head: Prof. Gianluca Bontempi.
- 10 researchers (2 prof, 4 postDoc, 4 PhD students), 4 graduate students.
- Research topics : Bioinformatics, Classification, Regression, Time series prediction, Sensor networks.
- Website : http://www.ulb.ac.be/di/mlg.
- Scientific collaborations inside ULB : IRIDIA (Sciences Appliquées), Physiologie Molculaire de la Cellule (IBMM), Conformation des Macromolcules Biologiques et Bioinformatique (IBMM), CENOLI (Sciences), Functional Genomics Unit (Institut Jules Bordet), Service d'Anesthesie (Erasme).
- Scientific collaborations outside ULB : UCL Machine Learning Group (B), Politecnico di Milano (I), Universitá del Sannio (I), George Mason University (US).

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Outline

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- Breast Cancer
- Traditional Approach for Prognostication
- Gene Expression Profiling Approach for Prognostication
- Isolation Content and Conte
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- Local Prognostic Gene Signatures

Onclusions

Introduction

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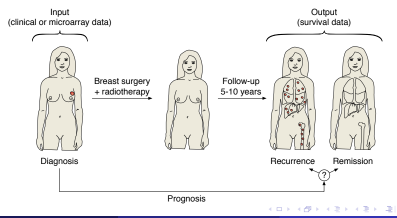
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Breast Cancer

• Huge public health issue:

- One of the most diagnosed malignancy in the Western World.
- ▶ 1 out of 9 women will develop a breast cancer during her lifetime.

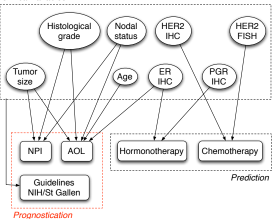
Prognostication:



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Traditional Approach for Prognostication

Clinical variables



- NPI = 0.2 × tumor size + grade + nodal status
- AOL uses a life table technique considering age, nodal status, tumor size, and ER.

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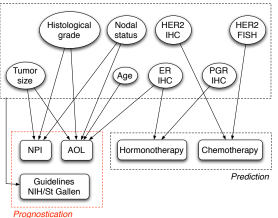
Problem: Prognostic clinical models predict numerous low-risk patients with early breast cancer (nodal status = 0) as high-risk.

Overtreatment

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Traditional Approach for Prognostication

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Overtreatment

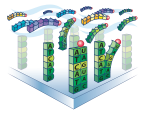
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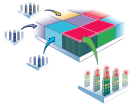
Microarray chip

Hybridization

Detection



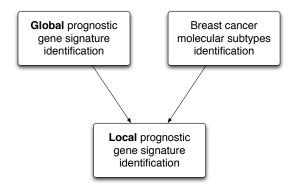




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Gene Expression Profiling Approach for Prognostication

- Improvement of breast cancer prognostication by using *machine learning* techniques to analyze microarray and survival data.
- Our methodology is composed of three main parts:

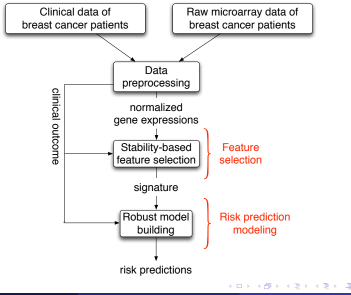


Global Prognostic Gene Signatures

- Objective: Identification of prognostic gene signatures without taking into account the presence of molecular subtypes.
- The idea is to identify prognostic gene signatures and their corresponding risk prediction models exhibiting the following characteristics:
 - Good performance with independent data.
 - Interpretable from a biological point of view.
 - Useable with data generated by different microarray platforms and/or normalization techniques.

Global Prognostic Gene Signature Identification

Methodology



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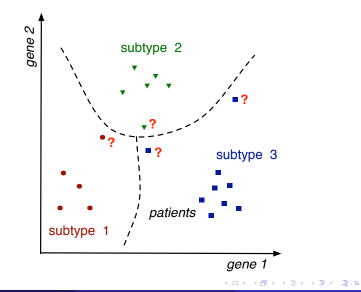
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Breast Cancer Molecular Subtypes

- Objective: Robust identification of breast cancer molecular subtypes along with estimation of classification uncertainty.
- The idea is to extent the study of Perou *et al.* by building a clustering model exhibiting the following characteristics:
 - Estimation of classification uncertainty.
 - Good performance in independent dataset.
 - Useable with data generated by different microarray platforms and/or normalization techniques.

Molecular Subtypes Identification

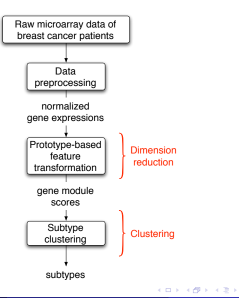
Classification uncertainty



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Molecular Subtypes Identification

Methodology



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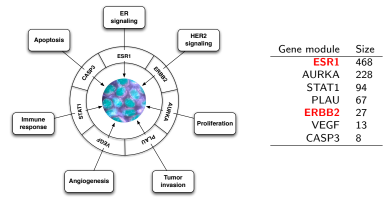
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- Aim: To reduce the dimensionality of microarray data while keeping information relevant for subtypes identification.
- We used a prototype-based feature transformation
 [Desmedt et al., 2008, Haibe-Kains, 2009] to identify sets of genes
 specifically co-expressed to key biological processes in breast cancer.
- The method is composed of 2 main steps:
 - Prototype-based clustering: Selection of prototypes and identification of gene modules.
 - Summarization: Each cluster is summarized by a single feature called gene module score (weighted average of genes in cluster).

 We applied this method with 7 prototypes [Hanahan and Weinberg, 2000] in two large datasets (NKI & VDX) [Desmedt et al., 2008].



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Subtype Clustering

- Aim: To develop of a robust clustering model able to estimate classification uncertainty.
- Low dimensional input space defined by ESR1 and ERBB2 module scores (see [Kapp et al., 2006]).
- We developed a model-based clustering:
 - Probability to belong to subtype j:

$$\Pr(j|x_i) = \frac{\pi_j \mathcal{N}(x_i; \mu_j, \Sigma_j)}{\sum_{l=1}^m \pi_l \mathcal{N}(x_i; \mu_l, \Sigma_l)}$$

where *m* is the number of Gaussians, π_j is the prior probability of x_i to be generated by the j^{th} Gaussian $\mathcal{N}(x_i; \mu_j, \Sigma_j)$

- Hyperparameters:
 - * Number of clusters (Gaussians): BIC [Schwarz, 1978].
 - Mean μ_j and covariance matrices Σ_j: EM algorithm [Dempster et al., 1977].

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• Pros

- The low dimensional space increases the model robustness (low feature-to-sample ratio).
- The low dimensional space facilitates the representation of the results and their interpretation.
- Estimation of classification uncertainty + ability to perform soft partitioning of the data.
- Easy to use this clustering model to predict the subtype of the tumor of a new patient (unlike hierarchical clustering).

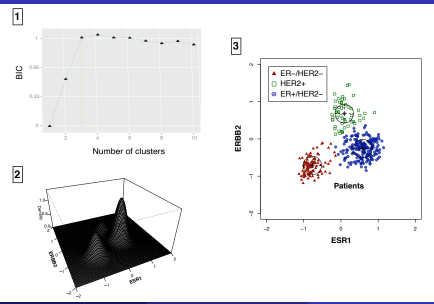
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 Although the use of ESR1 and ERBB2 module scores is beneficial (see above), it prevents us to find any new subtypes.

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- Training set: 344 patients with early breast cancer (VDX).
- Validation set:
 - Clustering: 17 independent datasets (> 3400 patients).
 - Prognosis: 745 untreated patients with early breast cancer from 5 datasets (NKI, TBG, UPP, UNT and MAINZ).

Subtype Clustering Model Training set (VDX)

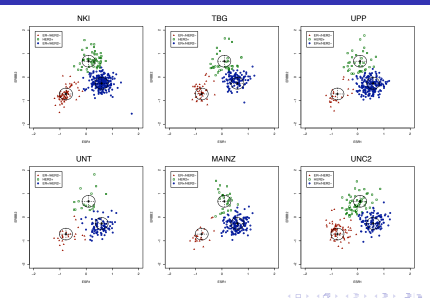


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Subtype Clustering Model

Independent datasets

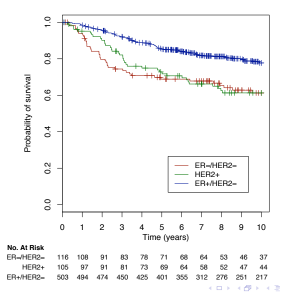


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Subtype Clustering Prognosis of early breast cancer patients



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Local Prognostic Gene Signatures

Local Prognostic Gene Signature Identification

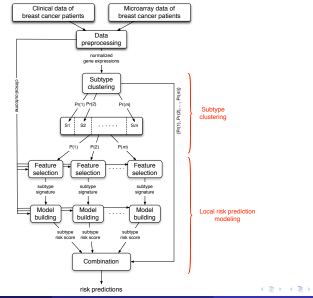
- Objective: Identification of prognostic gene signatures taking into account the presence of molecular subtypes.
- [Wang et al., 2005] conducted a study similar to [van't Veer et al., 2002] but they took into account the molecular heterogeneity of breast cancer wrt ER status (first *local signatures*).
- → Prognostic gene signature: GENE76.
 - Open issues:
 - The local models are developed for two subgroups (ER- and ER+) only, without considering the heterogeneity of the HER2+ subgroup.
 - Hard partitioning of the population of breast cancer patients.
 - The local model for ER- subgroup was trained on few samples and yielded poor performance in validation studies [Foekens et al., 2006, Desmedt et al., 2007].

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Local Prognostic Gene Signature Identification

Methodology



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Aim: To develop a prognostic model taking into account breast cancer molecular subtypes

• We adopted a *divide-and-conquer* strategy, through a *modular modeling* approach, and developed a Local Model Network (*fuzzy*) [Johansen and Foss, 1993] for prognostication:

$$r = \sum_{j=1}^{m} \rho_j(x, \theta_j) h_j(x, \alpha_j)$$

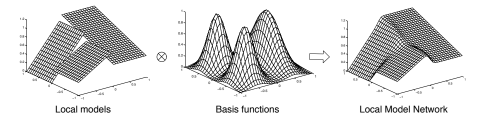
where x and r stand for the gene expressions and patients' risk respectively, $\rho_j(x, \theta_j)$ is the j^{th} basis function of parameters θ_j , $h_j(x, \alpha_j)$ is the j^{th} local risk prediction model of parameters α_j .

• The ρ_j are constrained to satisfy

$$\sum_{j=1}^m \rho_j(x,\theta_j) = 1$$

Local Model Network

• Example of LMN:



• The *j*th basis function is defined as the probability to belong to the *j*th breast cancer molecular subtype (*subtype clustering*):

$$\rho_j(x) = \frac{\pi_j \mathcal{N}(x; \mu_j, \Sigma_j)}{\sum_{l=1}^m \pi_l \mathcal{N}(x; \mu_l, \Sigma_l)}$$

• The *j*th local model is defined by our robust model:

$$h_j(x) = \sum_{i=1}^{k(j)} \beta'_i x_i$$

where signature size k now depends on subtype j (*local feature selection*).

- In order to take advantage of the subtypes identification, we adapted the stability-based feature ranking to identify the genes prognostic in specific subtypes.
- We introduced a weighted version of the concordance index:

$$C_{wted}(x_i, y, \rho_j) = \frac{\sum_{k,l \in \Omega} w_{kl} \mathbf{1}\{x_{ki} > x_{li}\}}{\sum_{k,l \in \Omega} w_{kl}}$$

where x_i are the expressions of gene *i*, *y* are the survival data and $w_{kl} = \rho_j(x_k)\rho_j(x_l)$ is the weight for the pair of *comparable* patients $\{k, l\}$ with $\rho_j(x)$ being the probability for a patient's tumor *x* to belong to the subtype *j*.

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Pros & Cons

Pros

- Use of robust linear risk prediction models to perform nonlinear modeling.
- Easy interpretation for doctors (breast cancer molecular subtypes + corresponding risk predictions).
- Potentially more biological insights since the molecular heterogeneity of breast cancer is taken into account.

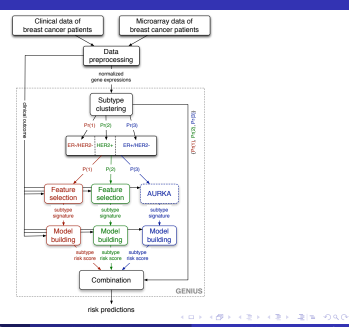
Local Prognostic Gene Signature Identification **GENIUS**

• GENIUS = Gene Expression progNostic Index Using Subtypes.

- Aim: Refining breast cancer prognosis according to molecular subtypes.
- GENIUS refers to:
 - The local prognostic gene signatures identified through local stability-based feature ranking.
 - The risk prediction model using robust model building and Local Model Network.
- Training set:
 - Input data: 22,283 (1050 after filtering) gene expressions of 344 untreated patients with early breast cancer (VDX).
 - Output data: Survival data.
- Validation set: 745 untreated patients with early breast cancer from 5 datasets (NKI, TBG, UPP, UNT and MAINZ).

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- Local prognostic gene signatures identified for the ER-/HER2- and HER2+ subtypes involved different genes but are enriched in genes related to immune response.
- GENIUS yielded significantly better performance than all the state-of-the-art prognostic gene signatures in the global population. GENIUS was not significantly superior in all subtypes but it was the only signature that performed well whatever the subtype.
- GENIUS significantly outperformed prognostic clinical models in the global population. Its superiority almost reached significance in all the subtypes.

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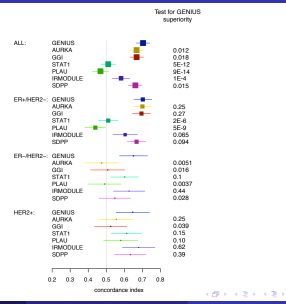
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Performance comparison with prognostic gene signatures



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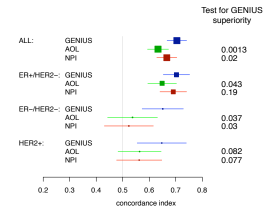


- Local prognostic gene signatures identified for the ER-/HER2- and HER2+ subtypes involved different genes but shared genes related to immune response.
- GENIUS yielded significantly better performance than all the state-of-the-art prognostic gene signatures in the global population. GENIUS was not significantly superior in all subtypes but it was the only signature that performed well whatever the subtype.
- GENIUS significantly outperformed prognostic clinical models in the global population. Its superiority did not reach significance in all the subtypes.

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Performance comparison with prognostic clinical models



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Conclusions

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Biological Insights

- From global prognostic gene signatures we highlighted the importance of proliferation when robustly quantified through gene expression profiling [Sotiriou et al., 2006, Wirapati et al., 2008, Desmedt et al., 2008].
- From local prognostic gene signatures we showed that the prognostic factors dramatically depend on the molecular subtypes, i.e. proliferation for ER+/HER2-, immune response for ER-/HER2- and HER2+ and angiogenesis for HER2+
 [Wirapati et al., 2008, Desmedt et al., 2008, Haibe-Kains, 2009].
- Since the ER+/HER2- subtype is the most common one (≈ 60% of the patients), we showed that the prognostic ability of most global gene signatures (e.g. GENE70, GENE76 or GGI) is driven by proliferation-related genes
 [Wirapati et al., 2008, Desmedt et al., 2008, Haibe-Kains et al., 2008b].

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- These findings suggested that a local modeling strategy taking into account for the presence of molecular subtypes might improve breast cancer prognosis.
- We showed that GENIUS yielded promising performance in independent datasets.
- The next step is to compare this fuzzy approach with a deterministic one to raise confidence in this local modeling strategy.



Improving the clinical value of tumor grading

<u>ManQuant</u> Dx[™] <u>Genomic Grade</u> is the cornerstone of the MapQuant Dx[™] assay series. It is the very first, microarray-based and clinically-validated, molecular diagnostic test to accurately measure tumor grade, a consensus indicator of tumor proliferation, risk of metastasis and response to chemotherapy.

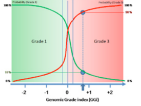
Resolving histological grading uncertainty

Tumor grade is a decision factor in most national & international guidelines to breast cancer treatment. It is generally recommended to treat high-grade "grade 3" breast carcinoma with

chemotherapy because they are chemosensitive and will often recur otherwise. By contrast, most low-grade "grade 1" tumors should not be treated with chemotherapy because they have a good prognosis and often are chemo-insensitive.

A critical clinical issue is how to treat the 50% of breast cancers tested today as intermediate grade?

The MapQuant Dx[™] Genomic Grade test now allows to resolve more than 80% of these uncertain "grade 2" tumors into "grade 1" or "grade 3" tumors, potentially sparing useless chemotherapy treatments to tens of thousands patients a year.



Clinical utility of the Genomic Grade index

CORPORATE

WEBSITE

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PRODUCTS

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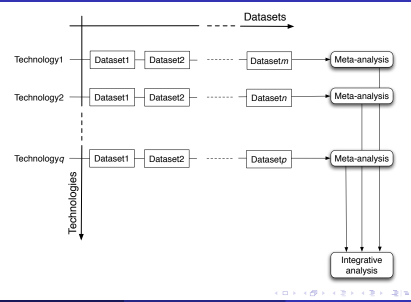
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Software

- survcomp R package available from CRAN:
 - Implementation of 6 performance criteria:
 - Hazard ratio
 - 2 D index
 - Oncordance index
 - Time-dependent ROC curve
 - Oross-validated partial likelihood
 - 6 Brier score
 - Implementation of 2 statistical tests for performance comparison:
 - ★ Paired student t test for 1, 2, and 3
 - ★ Wilcoxon signed-rank test for 4, 5, and 6.
- SWEAVE code: The LATEX and R codes performing the whole analysis of the microarray and survival data are publicly available for [Haibe-Kains et al., 2008a, Haibe-Kains et al., 2008b].

Future of Bioinformatics

Integrative bioinformatics



Thank you for your attention.

This presentation is available from http:

//www.ulb.ac.be/di/map/bhaibeka/papers/haibekains2009genius_talk_ki.pdf.

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Appendix

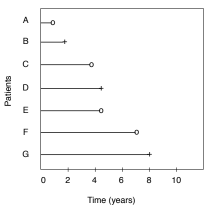
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• Retrospective observation plan (data retrieved from hospital DB).



Censoring

Survival data

Patient id	Time (years)	Event	
A	1	1	
В	2	0	
C	4	1	
D	5	0	
E	5	1	
F	7	1	
G	8	0	

Survival analysis

Censoring in survival data requires specific methods to deal with.

Survival Distributions and Performance

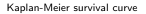
- Time of event occurrence are realization of a random variable **t**.
- Survival distributions:

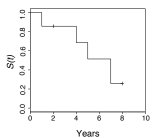
$$S(t) = 1 - \Pr\{\mathbf{t} \le t\}.$$

$$h(t) = \lim_{\Delta t \to 0} \frac{\Pr\{\mathbf{t} \le t\}}{\Delta t}$$

- Influence of covariates x:
 - Proportional hazards model (Cox): $h_i(t) = \lambda_0(t) \exp(\beta x_i)$

unspecified baseline



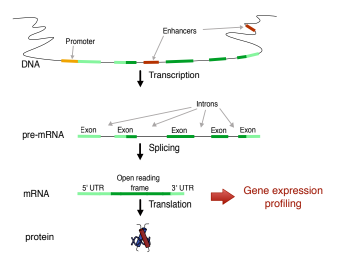


where x_i is the gene expression profile of patient *i* in the thesis.

risk

- Performance:
 - Concordance index: Estimate of the probability that, for a pair of randomly chosen comparable patients, the patient with the higher risk prediction will experience an event before the lower risk patient.
 - → C-index $\in [0, 1]$ (higher is better in the thesis).

Central Dogma of Molecular Biology



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Microarray Data

- Few samples (dozens to hundreds patients).
 - Microarray technology is expensive.
 - Frozen tumor samples are rare (biobank).
- Numerous gene expressions are measured (tens of thousands genes).
 - ► The recent microarray chips cover the whole genome (≈ 50,000 probes representing 30,000 "known genes").
- High feature-to-sample ratio (curse of dimensionality requiring feature selection).
- Microarray is a complex technology.
- High level of noise in the measurements.
 - Biology is complex (e.g. gene co-expressions due to pathways).

Redundancy.

• Microarray data matrix

DATA	<i>x</i> ₁	<i>x</i> ₂		хp
patient ₁	1.3	3.2		7
patient ₂	2.7	1.2		2.3
:	:	:	:	:
patient _n	5.4	1.4		9.1

Datasets

Dataset	Technology	Survival	Treatment Patients		Probes
VDX	Affymetrix	YES	untreated 344		22,283
NKI	Agilent	YES	heterogeneous	345	24,481
STNO2	cDNA Stanford	YES	heterogeneous	122	7,787
NCI	cDNA NCI	YES	heterogeneous	99	6,878
MGH	Arcturus	YES	hormono	60	11,421
MSK	Affymetrix	YES	heterogeneous	99	22,283
UPP	Affymetrix	YES	heterogeneous	251	22,283
STK	Affymetrix	YES	heterogeneous	159	22,283
UNT	Affymetrix	YES	untreated	137	22,283
UNC2	Agilent	YES	heterogeneous	248	21,495
DUKE	Affymetrix	YES	heterogeneous	171	12,625
CAL	Affymetrix	YES	heterogeneous	118	22,283
TBG	Affymetrix	YES	untreated	198	22,283
NCH	Agilent	YES	heterogeneous	135	17,086
DUKE2	Affymetrix	NO	chemo	160	61,359
MAINZ	Affymetrix	YES	untreated	200	22,283
TAM	Affymetrix	YES	hormono	354	44,928
TAM2	Aminolink	YES	hormono	155	21,332
LUND2	Swegene	NO	hormono	105	27,648
LUND	Swegene	NO	heterogeneous	143	26,824
MUG	Operon	NO	NA 152		16,783

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State-of-the-art: [van't Veer et al., 2002]

• Study of [van't Veer et al., 2002]:

- Training set:
 - ★ Input data: 24,481 (5000 after filtering) gene expressions of 78 untreated patients with early breast cancer.
 - Output data: Dichotomization of survival data (5 years).
- Feature selection: Feature ranking (correlation).
- Model building: Nearest centroid classifier.
- Hyperparameter tuning: Supervised tuning of the signature size (cross-validation).
- Validation set:
 - ★ Internal: Cross-validation !!!
 - ★ External: 295 treated and untreated patients with breast cancer at various stages [van de Vijver et al., 2002] (NKI) !!!

State-of-the-art: [Perou et al., 2000, Sorlie et al., 2001]

- Study of [Perou et al., 2000, Sorlie et al., 2001]:
 - Training set:
 - Input data: 8,102 gene expressions of 84 patients with breast cancer at various stages.
 - Feature selection: Feature ranking based on variance (*intrinsic gene* list).
 - Model building: Hierarchical clustering.
 - Hyperparameter tuning: Number of clusters is selected based on visualization assessment !!!
- Validation:
 - Internal: None !!!
 - External: 38 new patients in [Sorlie et al., 2003] + datasets from [van't Veer et al., 2002] and [West et al., 2001] !!!

State-of-the-art: [Wang et al., 2005]

- Study of [Wang et al., 2005]:
 - Training set:
 - Input data: 22,283 (17,819 after filtering) gene expressions of 115 untreated patients with early breast cancer.
 - ★ Output data: "Full" survival data and dichotomization (5 years) !!!
 - Feature selection: Feature ranking based on significance of univariate Cox's models (bootstrap) for ER- and ER+ separately.
 - Model building: Combination of univariate Cox's models.
 - Hyperparameter tuning: Signature size tuned by optimizing the sensitivity and specificity in the training set.
 - Validation set:
 - * Internal: 171 patients (single "random" split) !!!
 - * External: 180 untreated patients with early breast cancer.

Feature Selection

- Desirable properties of feature selection:
 - Computation efficiency (thousand of features).
 - Robustness to overfitting (not dataset dependent).
 - Intuitive tuning of (few) hyperparameters.
- For microarray data, **feature ranking** was shown to be efficient [Wessels et al., 2005].
- For prognostication, we used the **concordance index** [Harrell et al., 1996] as scoring function:

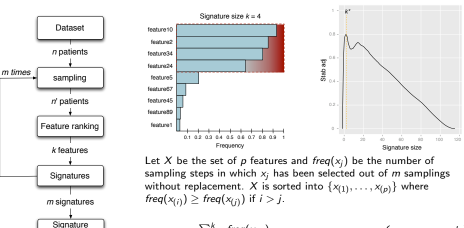
$$C ext{-index}(x_i, y) = rac{\sum_{k,l \in \Omega} \mathbbm{1}\{x_{ki} > x_{li}\}}{|\Omega|}$$

where x_i are the expressions of gene *i*, *y* are the survival data and Ω is the set of all the pairs of *comparable* patients $\{k, l\}$.

• Signature size?

Stability-Based Feature Ranking

Aim: To select a relevant and stable (low variance) set of features



$$Stab(k) = \frac{\sum_{l=1}^{k} freq(x_{(l)})}{k m} \qquad Stab_{adj}(k) = \max\left\{0, Stab(k) - \frac{k}{p}\right\}$$

$$k^* = \operatorname*{argmax}_k Stab_{adj}(k)$$

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stability

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Stability-Based Feature Ranking Pros & Cons

Pros & Cons

Pros

- Computational scalability.
- Reduction in risk of overfitting (low variance).
- Easy and intuitive tuning of the hyperparameter.

Cons

 Potential risk of bias since feature ranking is unable to deal with redundancy and complementarity of the features [Meyer, 2008].

This method was first applied in a study related to tamoxifen resistance in breast cancer [Loi et al., 2008].

- Risk prediction models can be:
 - Linear or non-linear.
 - Univariate or multivariate.
- Intrinsic nature of microarray data evokes the risk of overfitting of non-linear multivariate prediction models.
- But univariate linear prediction models are not able to take into account for the multiple interactions underlying tumorigenesis.

➡ Trade-off?

- Since 2002, large comparative studies have been conducted to identify successful prediction models for class discovery and classification
- ... leaving aside risk prediction (survival models) in breast cancer.

Comparative Study of Risk Prediction Models

- We compared numerous linear risk prediction models in the context of breast cancer prognostication using microarray and survival data [Haibe-Kains et al., 2008b].
 - "Complex" methods, although promising in the training set, yielded poor performance in independent sets (overfitting).
 - The loss of interpretability deriving from the use of overcomplex methods may be not sufficiently counterbalanced by significantly better performance.
- The few nonlinear risk prediction models (e.g. survival nnet [Eleuteri et al., 2003] and cSVM [van Belle et al., 2007]) developed in the field also yielded poor performance (data not shown).

• So we focussed on linear multivariate linear risk prediction models and relied on a simple additive combination scheme [Kittler et al., 1998] and equal weights linear regression [Wainer, 1976, Green, 1977]:

$$r = \sum_{i=1}^{k} \beta'_i x_i$$

where *r* is the *risk* of a patient, x_i are the gene expressions, and $\beta'_i = \begin{cases} -\frac{1}{k} & \text{if } x_i \text{ is positively correlated with survival} \\ +\frac{1}{k} & \text{otherwise} \end{cases}$

- The use of equal weights (coefficients)
 - reduces the risk of overfitting,
 - ensures the coefficients estimation to not be influenced by outliers,
 - and causes only modest expected loss in accuracy.

Robust Model Buidling Pros & Cons

Pros & Cons

- Pros
 - Additive models are multivariate models with low variance.
 - Less sensitive to redundancy that traditional multivariate models (collinearity).
 - Low computational cost (especially in combination with feature ranking).
 - Equal weights regression (i.e. signed average in our case) facilitates the computation of risk predictions in different microarray platforms with missing probes.

Cons

Unable to deal with complementarity of features.

This method was used in the study related to **GGI** = **G**ene expression **G**rade Index [Sotiriou et al., 2006].

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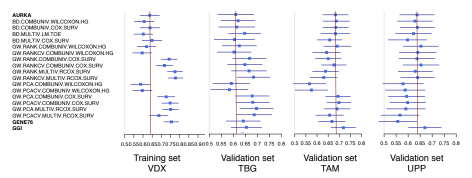
Comparative Study of Risk Prediction Models

- We compared numerous linear risk prediction models in the context of breast cancer prognostication using microarray and survival data [Haibe-Kains et al., 2008b].
- Risk prediction models:

	Genotype	Dim. reduction	Structure	Learning algo.	Phenotype
1	AURKA				
2	BD		COMBUNIV	WILCOXON	HG
3	BD		COMBUNIV	COX	SURV
4	BD		MULTIV	LM	TOE
5	BD		MULTIV	COX	SURV
6	GW	RANK(CV)	COMBUNIV	WILCOXON	HG
7	GW	RANK(CV)	COMBUNIV	COX	SURV
8	GW	RANK(CV)	MULTIV	RCOX	SURV
9	GW	PCA(CV)	COMBUNIV	WILCOXON	HG
10	GW	PCA(CV)	COMBUNIV	COX	SURV
11	GW	PCA(CV)	MULTIV	RCOX	SURV
12		. ,		GENE76	
13				GGI	

Comparative Study of Risk Prediction Models $_{\mbox{\scriptsize Results}}$

• Forestplot of the concordance index for each method in the training set and the three validation sets:



- "Complex" methods, although promising in the training set, yielded poor performance in independent sets (overfitting).
- The loss of interpretability deriving from the use of overcomplex methods may be not sufficiently counterbalanced by significantly better performance.

Global Prognostic Gene Signature Identification Biological validation

- **GGI** = **G**ene expression **G**rade Index.
 - Aim: Understanding the molecular basis of histological grade to improve prognosis.
- GGI refers to:
 - The signature of 128 probes selected through feature ranking with fixed threshold.
 - The robust model.
- Training set:
 - Input data: 22,283 gene expressions of 64 ER+ tamoxifen treated patients with breast cancer at various stages.
 - Output data: Binary class defined by histological grade 1 or 3.
- Validation set: 745 untreated patients with early breast cancer from 5 datasets (NKI, TBG, UPP, UNT and MAINZ).

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GGI Results

• Histological grade:

- HG1 and HG3 tumors have distinct gene expression profiles characterized by proliferation-related genes, HG2 tumors being heterogeneous.
- GGI is strongly predictive of HG1 vs HG3.

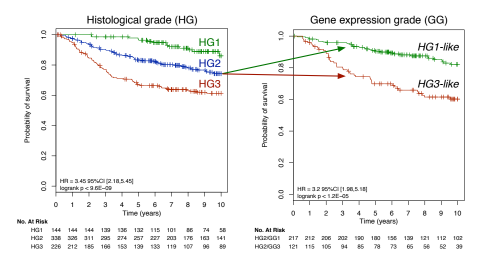
• Prognosis:

- Strong prognostic value: GGI is strongly associated with distant metastasis free survival (DMFS). The three-category histological grading system could be replaced with a two-catehory one using GG to improve prognosis.
- Proliferation and modeling: We showed in [Haibe-Kains et al., 2008b] that GGI is the only risk prediction model to outperform the single proliferation gene AURKA.
- Similar performance compared state-of-the-art gene signatures: GGI is competitive with GENE70 [van't Veer et al., 2002] and GENE76 [Wang et al., 2005].

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GGI Prognosis



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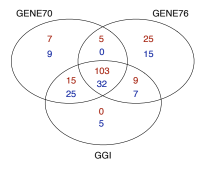
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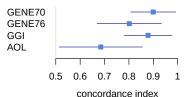
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• Concordance in predictions:



• Similar performance:



	Test difference	
GENE70 vs GENE76	0.15	
GENE70 vs GGI	0.53	
GENE76 vs GGI	0.22	

	Test superiority	
GENE70 vs AOL	0.007	
GENE76 vs AOL	0.13	
GGI vs AOL	0.015	

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- GGI is simple:
 - From a statistical point of view: robust model building.
 - From a biological point of view: GGI signature is mainly composed of proliferation-related genes.
- GGI yielded similar performance than gene signatures including genes related to many biological processes (GENE70 and GENE76).
 - Actually, we showed in [Wirapati et al., 2008] that the prognostic value of these signatures is mainly driven by the proliferation-related genes.
- The studies related to GGI challenged the use of risk prediction models of high statistical and biological complexity [Sotiriou et al., 2006, Haibe-Kains et al., 2008b, Desmedt et al., 2008].

EL OQA

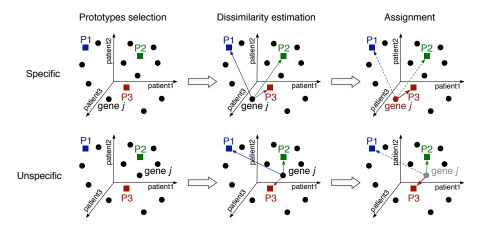
Molecular Subtypes Identification State-of-the-Art

• State-of-the-art:

- [Perou et al., 2000, Sorlie et al., 2001] were the first to study the molecular heterogeneity of breast cancer in order to build a subtype classifier.
- Open Issues:
 - * Lack of estimation of classification uncertainty.
 - The use of large number of genes led to a clustering model prone to overfitting (low robustness).
 - * The (number of) clusters were subjectively selected.
 - Lack of existing statistics to evaluate the robustness of the clustering model in independent datasets.
 - Association with survival tested in heterogeneous populations of breast cancer patients.
- [Kapp et al., 2006] confirmed the importance of ER and HER2 signaling pathways.

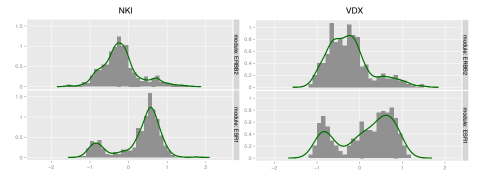
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Prototype-Based Clustering



-

Mixture of Gaussians?



-

Clustering Performance

- Performance assessment of clustering is a difficult task since the "truth" is hidden.
- In [Tibshirani and Walther, 2005], the authors introduced a new framework viewing clustering as a supervised learning problem in which the "true" class labels have to be estimated.
- Let X and Y be the training and validation sets, respectively.
- Let $C_X(Y)$ denote the use, in the dataset Y, of a clustering model C fitted on dataset X.
- Let $D[C_{\cdot}(.)]$ be the co-membership matrix of the clustering $C_{\cdot}(.)$.
- Idea:

 - 2 Cluster $X \rightarrow C_X(X)$.
 - **3** Cluster Y using $C_X \rightarrow C_X(Y)$.
 - Compare C_X(Y) and C_Y(Y) through the co-membership matrix D
 → prediction strength.

Subtype Clustering vs Perou's Method

Prediction strength in independent datasets (higher is better)

Dataset	Subtype clustering	Perou's method				
Dataset	3 clusters	2 clusters	3 clusters	4 clusters	5 clusters	
NKI	1.00	0.89	0.42	0.25	0.20	
TBG	0.83	0.92	0.39	0.24	0.22	
UPP	0.87	0.71	0.51	0.33	0.22	
UNT	0.89	0.78	0.55	0.36	0.00	
STNO2	0.69	0.86	0.44	0.28	0.25	
NCI	0.83	0.93	0.44	0.33	0.13	
STK	0.87	0.61	0.38	0.28	0.25	
MSK	0.96	0.77	0.66	0.20	0.00	
UNC2	0.87	0.81	0.57	0.31	0.00	
NCH	0.82	0.66	0.49	0.36	0.29	
DUKE	0.82	0.57	0.42	0.37	0.42	
DUKE2	0.64	0.92	0.63	0.47	0.00	
MAINZ	0.90	0.68	0.39	0.24	0.18	
CAL	0.95	0.84	0.41	0.31	0.00	
LUND2	0.87	0.92	0.51	0.17	0.17	
LUND	0.81	0.55	0.36	0.24	0.20	
MUG	0.49	0.50	0.33	0.27	0.23	
mean	0.83	0.76	0.47	0.30	0.16	
sd	0.12	0.14	0.10	0.07	0.12	

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- Personal webpage: http://www.ulb.ac.be/di/map/bhaibeka/
- Machine Learning Group: http://www.ulb.ac.be/di/mlg
- Functional Genomics Unit: http://www.bordet.be/en/services/medical/array/practical.htm
- Master in Bioinformatics at ULB and other belgian universities: http://www.bioinfomaster.ulb.ac.be/