Interactome-Transcriptome integration for predicting distant metastasis in breast cancer

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ABSTRACT

Motivation: High-throughput gene-expression profiling yields genomic signatures that allow the prediction of clinical conditions including patient outcome. However, these signatures have limitations, such as dependency on the training set, and worse, lack of generalization.

Results: We propose a novel algorithm called ITI (Interactome-Transcriptome Integration), to extract a genomic signature predicting distant metastasis in breast cancer by superimposition of large-scale protein-protein interaction data over a compendium of several gene-expression data sets. Training on two different compendia showed that the estrogen receptor-specific signatures obtained are more stable (11-35% stability), can be generalized on independent data, and performs better than previously published methods (53-74% accuracy).

Availability and Supplementary Information: The ITI algorithm source code and supplementary material from analysis are available under CeCILL from the ITI companion web site: http://bioinformatique.marseille.inserm.fr/iti.

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1 INTRODUCTION

The advent of post-genomic technologies provided the opportunity to potentially decipher the genomic origin of human diseases, including cancer. Thus, gene expression analysis using DNA microarrays allowed improving the classification and prognostication of several types of cancer, including breast cancer (Sørlie et al., 2001), van de Vijver et al. (2002). This approach can also help predict the metastatic recurrence and outcome (Wang et al., 2005). In breast cancer (BC), the current prognostic features poorly reflect the heterogeneous clinical outcome. The consequence is that many patients (70-80%) receive unnecessary adjuvant systemic chemotherapy. Genomic tools could provide an opportunity to refine prognosis and improve treatment strategy and lay down foundation of personalized medicine in BC.

Several studies have produced signatures linked to BC distant metastasis (Sotiriou et al., 2006). The Mammaprint 70-gene signature (van de Vijver et al., 2002) classified BC patients in either good or poor-prognosis groups. Wang et al (Wang et al., 2005) reported a 76-gene signature specific to Estrogen Receptor (ER) status (60 genes for ER+ patients, and 16 for ER-). These two signatures have an overlap of only 3 genes, which raised concerns about their reliability. Michiels and colleagues (Michiels et al., 2005) reanalyzed the van de Vijver dataset and concluded that the signatures obtained in such studies are unstable and dependent on the patients training set. From a classification standpoint, any gene classifier can be a good one as long as it performs and generalizes well (Dobbin et al., 2008). However, from either a scientific or clinical standpoint, both the content and stability of signatures are of primary importance, so as to decipher their molecular basis, to reinforce their robustness and widespread acceptance of their routine clinical use, and eventually to lead to new therapeutic targets.

Reasons for inherent instability of gene-based signatures have been previously enumerated (Fan et al., 2006, Bertucci et al., 2006). Besides experimental variability, variation in patient sampling and microarray platform bias, other reasons explain the lack of stability of signatures (Ein-Dor et al., 2006). Reasons best explaining this instability are (i) the curse of dimensionality, and (ii) the biological nature of gene expression measurements. The curse of dimensionality is well known of statisticians and is due to the inherent microarray data topology (too few samples for too many variables). The biological nature of instability is the following. Microarrays measure messenger RNA transcript abundance. To the extent that perturbations linked to a particular phenotype are reflected by changes in messenger RNA transcript levels, microarrays may be useful for measuring perturbations linked to a particular phenotype. Genes, however, are not independent but their products act in concert through protein-protein interaction network(s). Our hypothesis is that phenotypes such as cancer result from isolated and subtle molecular perturbations (changes in gene expression and/or mutations for example) in driver genes that may provoke expression changes of greater amplitude in downstream genes (Chuang et al., 2007).

Statistics for differential expression detect changes of greater amplitude and reveal only these downstream genes. Superimposing an interaction network to expression changes can detect driver genes associated with more subtle expression changes (Chuang et al., 2007). Factors cited previously may be more problematic for
markers for subtle changes in differential expression but we expect their effect to be attenuated by combination of several datasets. Such genes, used as biomarkers, have proved to be more robust in predicting distant metastasis of breast tumors profiled on heterogeneous platforms than genes detected without network information. Several network-based approaches have been published for microarray analysis. They include generating condition-dependent networks on differential expression, where no prior information on interaction data is used, which somewhat limits the biological relevance of the results (Gill et al., 2010). Co-clustering expression and graph data were proposed earlier by constructing a novel distance based on expression and network interactions (Hanisch et al., 2002). Support vector machines (SVM) in combination with spectral decomposition data denoising was proposed for analyzing transcriptional response in yeast (Rapaport et al., 2007). A network-based method was proposed to detect differentially-expressed subnetworks in existing Protein-Protein Interaction (PPI) data by local subnetwork aggregation (Chuang et al., 2007). Using a stricter statistical framework, an SVM variation for directly using interaction data within a classifier was applied to microarray classification (Zhu et al., 2009). These methods addressed the biological issues mentioned before. However, the data dimensionality issue was still not taken into account because training and testing were done on a single dataset with a relatively low number of samples. We propose here a multi-dataset re-implementation of the method proposed by Chuang et al. (2007) to integrate analysis of several gene expression datasets so as to extract subnetworks discriminating BC distant metastasis. We demonstrate the performance of our method, called Interactome-Transcriptome Integration (ITI) on a large compendium of publicly available data. To avoid potential bias in subnetwork selection, we performed a stratified ten-fold cross-validation and combined the obtained networks. Validation was then done on two independent BC gene expression datasets (Desmedt et al., 2008, van de Vijver et al., 2002). Using this approach, we significantly increased the classification performance as compared to three previously published signatures while lowering the dependence of the signature on the training set. Independent classification on two studies by van de Vijver and coworkers (2002) and Wang and coworkers (2006) achieved 53 and 74% accuracy, respectively. We detail here our ITI algorithm and report statistical validation, patient classification results, as well as biological validation of the subnetworks thus defined.

2 METHODS

To detect protein complexes with subtle expression changes, we superimposed a large scale PPI map to a compendium of BC expression profiles. The strategy implemented in ITI consists in detecting interactome subsets (subnetworks) whose expression is significantly correlated with Distant Metastasis-Free Survival (DMFS) in several datasets simultaneously. These subnetworks are then validated subsequently by shuffling interaction data and gene expression data. To train and test the system, six public datasets were chosen according to the criteria section 2.2. Four analyses were performed (two different validation datasets held out for independent testing for study 1 (Desmedt’s dataset) and study 2 (van de Vijver’s dataset) and separate analysis according to positive or negative ER tumor status) to assess the impact of training data on the detected subnetworks and to understand their generalization capability. For each study, a ten-fold cross-validation was performed by carefully stratifying the training (90% of samples) and test sets. The aim of stratification was to properly balance each of the ten training sets to keep the same ER+/ER- and DMFS events proportions in each of the 10 iterations.

2.1 Protein-Protein Interaction data integration

The following interaction data sets were used: Human Protein Resource Database release 9 (HPRD Keshava Prasad et al., 2009), Molecular Interaction database (MINT, Ceol et al., 2010), INTAct (Aranda et al., 2010), Database of Interacting Proteins (DIP, Salwinski et al., 2004), and the human interactome generated in silico with the Cocite algorithm (Ramani et al., 2005). All data were downloaded the 8/9/2010, and parsed to remove self-interactions, duplicates and proteins marked as “unknown”. Self interactions were removed from the files as they are not quantified by the algorithm, and interaction maps were integrated by uniqueness of NCBI gene ID accession numbers. Annotations were homogenized within datasets for proper display within the system. Resulting interactions obtained by crossing all interaction data totaled a number of 70530 single interactions among a total of 13202 proteins.

2.2 Breast cancer compendium (BCC)

The public datasets (see Table 1) were selected and included in a Breast Cancer Compendium (BCC) on the basis of the following criteria: early breast cancer, availability of clinical information related to metastasis (event information and delay between the BC diagnosis, and the relapse or the last follow-up) and immunohistochemical ER status (ER+, ER-) and absence of post-operative adjuvant chemotherapy. A total of 930 tumors were retained for analysis from the initial pool of 1561 tumors through 6 datasets. Sampling size, platform types and ER status are detailed in Table 1. DMFS status was censored if follow-up was less than 5 years for all datasets.

Table 1. Datasets included in the breast cancer compendium. Two trainings (study 1 and 2) were performed using different combinations of training and testing data (on bold): On the study 1, Desmedt tumors were held out for independent testing, and training was done on the rest. Van de Vijver dataset was respectively held out for the study 2.

<table>
<thead>
<tr>
<th>Author(s)</th>
<th>GEO Accession</th>
<th>Platform</th>
<th>Samples</th>
<th>DMFS Status</th>
<th>ER/ER+</th>
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<tr>
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<td>Lai et al., 2008</td>
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<td>HG-U133A and B</td>
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<td>27/74</td>
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<td>Schmidt et al., 2008</td>
<td>GSE11121</td>
<td>HG-U133A</td>
<td>182/200</td>
<td>46/136</td>
<td>37/145</td>
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<tr>
<td>van de Vijver et al., 2002</td>
<td>N/A</td>
<td>Agilent Hu-manGenome</td>
<td>150/295</td>
<td>56/14</td>
<td>36/114</td>
</tr>
<tr>
<td>Wang et al., 2005</td>
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<td>HG-U133A</td>
<td>276/286</td>
<td>107/169</td>
<td>72/204</td>
</tr>
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<td>6 publicly available</td>
<td>4 distinct platforms</td>
<td>930/1561</td>
<td>307/623</td>
<td>246/684</td>
</tr>
</tbody>
</table>

Raw data sets were downloaded from National Center for Biotechnological Information (NCBI) Gene Expression Omnibus repository (Barrett et al., 2009) when available, and normalized using the GCRMA method from Bioconductor. The van de Vijver dataset was downloaded as supplementary material from the publication (van de Vijver et al., 2002). Datasets were collapsed from probe expression to gene expression as described in


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Reyal et al. (2005). When multiple probesets were available for a gene, we used the probeset having the highest median signal. Following this, ‘nx_at’ marked probes were removed. HG-U133A and HG-U133B were integrated as a virtual combined platform.

2.3 Dataset stratification, Imbricated ten-fold cross-validation and independent testing

To detect discriminative subnetwork while avoiding over-fitting, cross-validation was performed by building training/testing sets while taking into account the clinical and molecular status of the tumors. Hence, stratification was done to balance ER+/ER- and distant metastasis rate between training and testing sets, leading to ten randomly selected training sets. Preservation of both molecular and clinical status proportions in each dataset allowed increasing training and testing sets homogeneity and avoided molecular bias.

For each test/train set, subnetworks were detected with the ITI algorithm (see section 2.4) and validated by gene expression and PPI shuffling (section 2.5), yielding five subnetwork lists. The lists were combined into a single signature (section 2.6) whose discriminative power was tested on datasets held apart, as described section 2.7.

2.4 Interactome Transcriptome Integration - Constructing subnetworks

Each couple of training/testing set was searched for discriminative subnetworks whose average expression was linked to distant metastasis using the ITI algorithm. The latter is derived from the algorithm of (Chuang et al., 2007), with the added capability of detecting subnetworks on a compendium (Figure 1). ITI was implemented as a pipeline developed with open source interpreters Perl and Bash and statistical validation was implemented with Matlab Statistical Toolbox R2010b (The Mathworks (c) Natick, MA, USA). Subnetwork detection was parallelized and implemented on a Beowulf Cluster to reduce computing costs. Subnetworks visualization was obtained with the graph layout package GraphViz (AT&T Research, USA).

To detect discriminative subnetworks, correlations between clinical status and gene expression were computed for each dataset. Then, the interactome was exhaustively explored for discriminative regions (see Figure 1) by individually considering each node as a potential seed and aggregate recursively neighbors on the basis of a score measuring correlation of expression with DMFS status (see equation 1). Neighbors were aggregated until subnetwork score could not be improved above a certain threshold (improvement score threshold = 0.03). Then, the following node in interactome was processed. Parallelization was done by subdividing interactome over available datasets, as described section 2.4.

Validating subnetworks - filtering

To validate subnetworks statistically, two random distributions of score were drawn. The first random distribution assessed the significance of algorithm that extracts subnetworks. It was obtained by randomly selecting subnetworks, i.e. by randomly accepting whether a subset of the interactome is a subnetwork without taking gene expression data into account. The second distribution assessed statistical significance of the biological link expression-protein-protein interaction. It was obtained by shuffling clinical conditions. To keep random subnetworks comparable to detected subnetworks, their distribution of size was forced to match that of the selected subnetworks by Gaussian modeling. Next, the distributions of random subnetwork scores were modeled by mixture of two Gaussian distributions. Once obtained, these distributions were used to fix scores thresholds independently over all datasets at significance levels of p-values, and filtered statistically significant subnetworks. Shuffling random interactions to obtain a random interactome did not yield subnetworks at reasonable score levels, confirming the strong link between gene expression levels and protein-protein physical interaction(s). Finally, we kept only the subnetworks with a score higher than expected by chance both on subnetwork randomization (p<1.10^-4 on 2 datasets) and shuffling of expression (p<1.10^-4 on 2 datasets).

2.6 Constructing a common subnetwork signature for each training set

Using this filter, 10 subnetwork sets for each training tumor subset were generated. Next, these sets were combined by examining subnetworks pair by pair across datasets and combining them if overlap was larger than 50%. Using this method, clusters of overlapping subnetworks were built. Finally, a subnetwork list was constructed from the list of subnetwork clusters by
keeping only subnetworks appearing at least twice. For a given cluster, only the subnetwork with the highest score was kept. Final subnetwork sets size are detailed Table 2.

2.7 Tumor classification and distant metastasis prediction of ER+ and ER- tumors on two independent datasets

The subnetwork list obtained section 2.6 was used for independent classification using two different settings, namely study 1 (in which Desmedt’s data was held out) and 2 (in which van de Vijver’s dataset was held out). In each setting, training was performed separately on tumors from all datasets except the held out dataset, yielding 5 SVM models. Classification on the validation sets was done by majority vote (weighted by sample size for each dataset) on the 5 SVM models. A complete organization chart is presented in Figure 2.

To use subnetworks as unique SVM input variables, gene expression within a subnetwork was averaged over genes and used as a discriminative profile for both training and testing. Several SVM models were tested for increasing number of subnetworks. A final subnetwork list was retained by maximizing accuracy. Classification results (accuracy, true and false positives) are reported in section 3, along with a comparison with previously published classifiers.

Fig. 2: Complete data workflow. Interactome is assembled from multiple sources (see section 2.1). Gene expression datasets forming our Breast Cancer Compendium (BCC) assembled section 2.2 are pooled to form a training dataset. Ten groups were then formed on a 10% leave-out basis (section 2.3). Subnetworks were detected on human interactome on each training set using ITI (section 2.4) and validated twice by shuffling interactions and expression, as described section 2.5. Retained subnetworks were combined (section 2.6) to train a Support Vector Machine (section 2.7). Final set was then used as a set of markers for classifying independent data by majority vote on the 5 SVM models (section 2.7).

2.8 ITI on-line resource - Gene Ontology category enrichment

To detect pathways associated with BC distant metastasis, we computed enrichment of biological process gene ontology in each subnetwork detected by ITI using the ErmineJ program (Gillis et al., 2010) and the reference list of Biological Process from Gene Ontology (The Gene Ontology in 2010: extensions and refinements, 2010). ErmineJ provided corrected p-values for enrichment of ontological terms computed with hypergeometric distribution. These were systematically computed for all subnetworks to associate them to known molecular processes defined in the Gene Ontology.

The resulting data were organized in a dedicated on-line resource (http://bioinformatique.marseille.inserm.fr/iti). This resource describes subnetworks detected with ITI and gives a thorough description of the included genes. Subnetworks and gene lists are downloadable for further processing. Subnetworks p-values calculated according to random distributions described in section 2.5 were also included, along with combined Fisher scores (Hong et Breitling, 2008). Genes were annotated with direct NCBI EntrezGene links and links to other subnetworks are provided. To understand expression changes of genes included in subnetworks, color-coded gene graphs are provided, with correlation expression/DMFS status superimposed on subnetworks. The correlation score is provided for all datasets separately.

3 RESULTS

3.1 Establishment of two discriminative subnetwork sets (ER+ and ER-) from a joined compendium of 930 tumors

Two separate signatures were generated for ER+ and ER- BC subtypes for two studies. In study 1, Desmedt’s data (Desmedt et al., 2008) was held out, and in study 2, van de Vijver’s data (van de Vijver et al., 2002) was held out, as described in section 2.3. Thus, four sets of subnetworks were assessed (see Table 2).

Table 2. p-values thresholds and signature size for the four training configurations (study 1 = all BCC but Desmedt, study 2 = all BCC but van de Vijver). The optimal number of subnetworks for classification depends on the training set and is lower for ER+ tumors, which reflects a higher homogeneity.

The optimal signature size retained in Table 2 is the one that maximizes the average accuracy on the 10 training sets for each analysis. For the study 1, discriminative subnetworks had an average score of 0.49 (ER+) and 0.54 (ER-) confirming the high correlation of co-expression and proximity in the PPI network. Signature size was respectively of 6 (ER+) and 165 subnetworks (ER-). For the study 2, the ER+ signature yielded an optimal classification score on independent data for 14 subnetworks, and the ER- signature for 122 subnetworks. They correspond to lists of 175 (Study 1, ER+), 2310 (Study 1, ER-), 272 (Study 2, ER+), and 1481 (Study 2, ER-) genes, respectively, many genes being represented in several subnetworks. These numbers are larger than what is reported for other signatures. This suggests that we detected a large panel of genes significantly linked to distant metastasis, realistically reflecting both the biological footprint of metastasis and the scale of perturbations at the gene expression level. Redundancy of genes within subnetworks may be explained by the high connectivity of several hubs (for instance TP53), which makes them likely to be included in several subnetworks.
To assess the performance of signatures constructed with ITI, we tested. Performance was measured on the same tumors (Desmedt and van de Vijver datasets), separately on ER+ and ER- tumors. The classification methods from the respective original publications were used for each signature. For van de Vijver’s signature, distances to mean centroids from relapse and non-relapse groups are calculated (van de Vijver et al., 2002), and the 76-gene ER status-specific signature (Wang et al., 2005) were tested. For Wang’s signature, a relapse score is calculated for each patient by a linear combination of expression weighed by standardized Cox’s coefficients (Wang et al., 2005). Because the GGI and Mammaprint signatures are probe-specific, the tests were done with the probes present in the test dataset. Results and performance measurements are detailed in Table 3. They show that ITI generalization performance is vastly superior to previously published signatures. The GGI classification showed the highest accuracy on the [47%-68%] range, the 70-gene signature on the [41%-62%] range, and the 76-gene signatures on the [37%-63%] range.

ITI gave a better accuracy as compared to the Wang signature on Desmedt’s data (ER+); an accuracy of 74% (specificity of 92%) was obtained versus an accuracy of 60% (specificity of 56%) with the 76-gene signature. ITI gave superior results also on Desmedt’s ER- tumors with an accuracy of 54% (specificity of 65%) versus an accuracy of 38% (specificity of 41%) for the Wang signature. This held true for the Mammaprint 70-gene signature, which works mostly for van de Vijver patients. ITI showed an accuracy of 53% associated with a specificity of 90% on van de Vijver’s data (ER-) and an accuracy of 52% with a specificity of 65% on van de Vijver’s ER+ tumors. This performance is inferior to what was obtained on study 1 and may reflect a bias towards Affymetrix induced by the training compendium. The Mammaprint signature had a lower performance of 41% on ER+ and of 42% on ER- Desmedt tumors. Similarly, ITI showed performance superiority over the GGI for ER- patients. Overall, ITI was able to generalize better with a lower accuracy bound of 52%.

On a different comparison basis, Chuang and coworkers (2007) achieved 48.8% accuracy on van de Vijver samples with training on Wang samples and 55.8% reciprocally. Specific contributions of the interaction data or gene expression data are not quantified, since they are not easily separable in the current setting. However, Chuang and coworkers (2007) already demonstrated that an expression approach increased signature robustness, and several studies showed that gene expression meta-analysis also increased classification performance (Xu et al., 2005, Fishel et al., 2007).

We performed a survival analysis between good and poor prognosis groups in study 1 (ER+) (Figure 3). Log-rank test gave a p-value of 4.89x10^-3, suggesting good separation between the two groups. This is higher than p-values obtained with other signatures (Wang signature gave p=4.11x10^-3, and GGI gave p=1.34x10^-3). The Mammaprint signature was not able to separate Desmedt’s patients in significant groups. Even though ITI was not specifically designed to obtain good log rank score, it was able to separate patients with higher survival and patients with lower survival expectancy. An alternative could have been to compute subnetwork score directly on genes log-rank p-values.
Fig. 3. Kaplan-Meyer estimator of good-prognosis (lower risk of distant metastasis) and poor-prognosis groups (higher risk of distant metastasis) survival rates as defined by ITI, Mammaprint, Wang’s signature and GGI for the ER+ Desmedt dataset. ITI gave the lowest p-value of $4.89 \times 10^{-5}$ with a Log-rank test among all tested signatures.

3.3 Signatures obtained with ITI show a stability of 11.5-32.8% for different training sets

Wang and van de Vijver signatures have only 3 genes in common, which represent less than 5% of all the genes in the two signatures. We compared the two signatures obtained with ITI for ER+ and ER- samples with the Desmedt and van de Vijver tumors. A total of 937 genes were found in common between the Desmedt and the van de Vijver signature for ER- samples, and 46 genes between the Desmedt and van de Vijver signatures for ER+ samples. This represents an overlap of respectively 32.8% (ER-) and 11.5% (ER+). These relatively low values reflect the fact that datasets and platforms are biased. However, this is largely superior to the 3 common genes between the Wang and van de Vijver signatures. This overlap between subnetwork sets could probably be improved by using a larger training compendium.

3.4 Biology of the discriminative subnetwork set is meaningful

We examined the enriched annotations from the Gene Ontology biological process for the subnetworks obtained in study 1. Table 4 shows several enriched GO terms for both ER+ and ER- signatures. Ontology terms found in discriminative subnetworks are linked to regulatory processes disrupted in cancer (cell cycle, DNA damage checkpoint) and in metastasis (immune system, cell proliferation, focal adhesion, cell migration and cytoskeleton organization) in both ER+ and ER- tumors.

As an example, we describe here a subnetwork significantly associated with metastasis in study 1 (ER-) (subnetwork 6693, represented in Figure 4). Subnetwork 6693 contained genes with well known function in ER- BCs and metastasis, such as the tumor suppresor gene (TSG) TP53 and the tyrosine kinase receptors ERBB2 and EGFR.

The subnetwork contained also several cell cycle kinases and regulators (CDK2, CDKN1A, CDKN2A). NQO1, whose altered expression has been associated with various forms of cancer. PIN1 is present in the subnetwork, and was recently found to promote aggressiveness in BC. Insulin receptor was also present; its deregulated expression correlates with poor response to anti IGF-FR therapy in triple negative BC. It also contained several well known oncogenes and genes not previously linked to cancer, but which may be acting as BC driver genes.

![Fig. 4. Graphical representation of part of subnetwork 6693 (Study 1, ER-). This illustrates a discriminative subnetwork from the Sabatier and coworkers dataset. Nodes and edges correspond to genes encoding proteins and protein-protein interactions, respectively. Yellow and blue nodes denote an over-expression and an under-expression, respectively, among patients with distant metastasis compared with the other ones.](http://bioinformatics.oxfordjournals.org/)

<table>
<thead>
<tr>
<th>Gene Ontology</th>
<th>GO</th>
<th>Corrected p-value</th>
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<td>ER+</td>
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<td>mRNA cleavage</td>
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<td>Wnt receptor signalling pathway through beta-catenin</td>
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</table>

Table 4. Enriched Gene Ontology annotations of ER+ and ER- subnetworks. Several enriched ontologies for subnetworks extracted in study 1 (ER-) and study 1 (ER+) studies are related to cancer.
4 DISCUSSION

We conceived a network-based algorithm (Interactome-Transcriptome Integration) to identify prognostic genomic signatures generalizable over multiple and heterogeneous microarray datasets. This algorithm works in two steps: first it integrates data from a compendium of BC microarray datasets, and second it finds subnetworks, i.e. interacting gene complexes, whose expression discriminates two conditions of interest. Subnetworks are filtered by statistical validation.

We applied the ITI algorithm to the particularly important but still unresolved question of finding markers for BC distant metastasis for which a large body of public data is available.

Our approach illustrates the feasibility of integrating gene expression data compendia (930 BC tumor samples were integrated) and large scale PPI data; it represents a potential data mining tool for gene expression repositories. It features inclusion of prior data under the form of PPI interactions and clinical annotations.

We produced two ER status-specific signatures that were validated on independent datasets held out from training. Repeating the experiments for two datasets (Desmedt and van de Vijver) yielded higher classification performance than previously published classifiers in both cases [74% for Desmedt (ER+) and 53% for van de Vijver (ER+)]. Our subnetwork-based signatures reflect the large biological footprint of metastasis and is consequently larger than previously published signatures. The classifier obtained with ITI subnetworks was less sensitive to platform bias than previously published classifiers, since performance obtained was similar on the two training compendia. It also showed high specificity, which is critical to make a decision on avoiding unnecessary adjuvant systemic treatment.

The ITI algorithm is currently extended to incorporate other data types, including DNA copy number variation data [SNPs, Comparative Genomic Hybridization arrays (CGH) and DNA methylation profiles]. ITI capability to handle the curse of dimensionality makes it suitable to detect biomarkers yielded by deep sequencing analysis. In next versions, PPI interaction type will also be taken into account at the interactome integration and subnetwork aggregation steps. Also, classification performance is inherently tied to molecular subtypes and finer subtyping is necessary to render this technology suitable for clinical use. A significant increase in ER-classification was observed by separating early and lately relapsing patients (data not shown). Further clinical validation could be envisioned through a phase-2 clinical trial with customized microarrays for adjuvant chemotherapy treatment decision making.

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Conflict of interest: none declared.

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