Translating bioinformatics in oncology: guilt-by-profiling analysis and identification of KIF18B and CDCA3 as novel driver genes in carcinogenesis

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ABSTRACT

Motivation: Co-regulated genes are not identified in traditional microarray analyses, but may theoretically be closely functionally linked [guilt-by-association (GBA), guilt-by-profiling]. Thus, bioinformatics procedures for guilt-by-profiling/association analysis have yet to be applied to large-scale cancer biology.

We analyzed 2158 full cancer transcriptomes from 163 diverse cancer entities in regard of their similarity of gene expression, using Pearson’s correlation coefficient (CC). Subsequently, 428 highly co-regulated entities in regard of their similarity of gene expression, using Pearson’s correlation coefficient (CC). Subsequently, 428 highly co-regulated genes (CC ≥ 0.8) were clustered unsupervised to obtain small co-regulated networks. A major subnetwork containing 61 closely co-regulated genes showed highly significant enrichment of cancer bio-functions. All genes except kinesin family member 18B (KIF18B) and cell division cycle associated 3 (CDCA3) were of confirmed relevance for tumor biology. Therefore, we independently analyzed their differential regulation in multiple tumors and found severe deregulation in liver, breast, lung, ovarian and kidney cancers, thus proving our GBA hypothesis. Overexpression of KIF18B and CDCA3 in hepatoma cells and subsequent microarray analysis revealed significant deregulation of central cell cycle regulatory genes. Consistently, RT-PCR and proliferation assay confirmed the role of both genes in cell cycle progression.

Finally, the prognostic significance of the identified KIF18B- and CDCA3-dependent predictors ($\overline{P} = 0.01$, $P = 0.04$) was demonstrated in three independent HCC cohorts and several other tumors.

In summary, we proved the efficacy of large-scale guilt-by-profiling/association strategies in oncology. We identified two novel oncogenes and functionally characterized them. The strong prognostic importance of downstream predictors for HCC and many other tumors indicates the clinical relevance of our findings.

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1 INTRODUCTION

Cancer is one of the leading causes of death (Jemal et al., 2009). Despite evident improvements in diagnostic procedures and the development of novel and effective therapies, the prognosis of disease remains dismal for many patients. A thorough understanding of the molecular basis of cancer development is critical to improve therapeutic options for a variety of cancers (Baehner et al., 2011). However, the identification of genes causally associated with complex diseases such as cancer remains challenging and therefore constitutes a major barrier in advancing our limited mechanistic understanding of the disease (Ioannidis, 2010). Simultaneously, increasing awareness of the fact that high-throughput gene expression analysis may reveal valuable insights into cancer biology led to the establishment of a variety of cancer microarray datasets and large microarray dataset repositories (Barrett et al., 2013; Davis et al., 2007; Parkinson et al., 2011; Sherlock et al., 2001). However, once published, these datasets are most often neglected as regards to further analysis. In fact, meta-analyses of this vast body of data are rarely attempted (Cahann et al., 2007; Krupp et al., 2011). The reasons are manifold, ranging from the difficulty of integrating various technical platforms to the absence of bioinformatics knowledge on the part of primarily molecular oriented scientists, and the need for advanced statistics (Ioannidis, 2010). Finally, these datasets could be subject to alternative analysis strategies to obtain additional information not yielded by ‘standard’ analysis protocols.

The concept of transferring gene function annotation from one gene to another based on the profile of their characteristics (guilt-by-profiling’, guilt-by-association (GBA)) has been frequently described. New biological relationships have been successfully identified in GBA studies, including correlated gene expression (Wu et al., 2002), correlated phylogenetic profiles (Dute et al., 2003) and membership within gene expression clusters (Hughes et al., 2000; Stolovitzky 2003).
Some guilt-by-profiling studies have exploited gene–gene relationships by transforming them into gene functional characteristics. This occurs when integrating gene expression profiling or functional annotations of the respective genes (Wu et al., 2002). Despite such proven success, no attempts have been made to include samples from multiple tumor entities or samples of both types of inference. The benefits of such a large-scale GBA approach are obvious because genes closely regulated across ≥2000 samples and various tumor entities may be active in a highly robust carcinogenic network. We therefore hypothesized that integrating multiple sources of biological information, such as expression profiling and functional annotation over thousands of tumor samples and entities, may provide novel insights into key regulatory changes in cancer development and help to identify highly causative genes.

Using this approach and proving its feasibility, we validated the use of bioinformatics GBA strategies in large-scale molecular oncology, identified two novel oncogenes, demonstrated their deregulation in several tumors, functionally characterized the novel oncogenes, identified corresponding downstream targets and showed the prognostic relevance of downstream predictors for hepatocellular carcinoma (HCC) as well as many other tumors (Fig. 1).

2 METHODS

2.1 Bioinformatics data analysis

The ‘GSE2109’ dataset was obtained from NCBI’s Gene Expression Omnibus 39 via the Bioconductor package ‘GEOquery’ 40. Further analysis was performed using R (http://r-project.org). According to the experimental description, signal values of the probe set were summarized using the Affymetrix Suite 5.0 (MASS) and normalized. After downloading and combining the data into a single expression set, the expression data were transformed for each array via the Z-score (Cheadle et al., 2003). Gene-centered information was obtained by summarizing and averaging the expressions of all gene-specific spots per array as described by the annotation GPL570 and documented in the Gene Expression Omnibus. Highly correlated gene expressions were detected by Pearson’s correlation coefficient (CC). Genes with a CC of $|CC| > 0.8$ were used for further analysis. For hierarchical clustering, distances between genes within the reduced dataset were calculated with Pearson’s CC and transformed through $CC = \frac{1}{1 + |CC|}$, to be used for hierarchical clustering. Complete clustering was applied to transformed distances. To estimate the ideal number of clusters, the KL divergence was used for hierarchical clustering. Complete clustering was applied to transformed distances between genes within the reduced dataset were calculated with Pearson’s CC and transformed through $CC = 1 - CC$. Analysis of CC was performed in C++ because of the enormous quantity of data, and for parallelization by Phthreads, which significantly accelerated the analysis. An optimal number of subclusters were identified using the C and KL indices (Charrad et al., 2010).

2.2 Analysis of gene expression correlation

The CC is a measure of the linear interdependence of the characteristics. We used Pearson’s CC to calculate correlating gene expression, which ranged between −1 and 1. Analysis of CC was performed in C++ because of the enormous quantity of data, and for parallelization by Phthreads, which significantly accelerated the analysis. An optimal number of subclusters were identified using the C and KL indices (Charrad et al., 2010).

2.3 Analysis of robustness of co-regulated genetic subclusters

Main data were reduced to the specific tissues of interest (liver, colon and breast). Robustness was calculated by applying the methods described above to the selected tissue-specific subsets. Genes within cluster #4 in the main data analysis were subsequently mapped to the novel tissue-specific clusters and analyzed for overlapping genes, genes with changing associations and genes not being co-regulated >0.8 in specific tissue.

2.4 Cell lines and vectors

The target gene kinesin family member 18B (KIF18B) and cell division cycle associated 3 (CDC3) sequences were obtained from the NCBI Gene. The annotated sequences have been cloned in a PUC57 vector. For overexpression in the mammalian cell line, a PCI vector carrying a cytomegalovirus (CMV) promoter was used (Promega). HUH7 cells were cultivated in advanced Dulbecco’s modified Eagle medium (DMEM). The cells were seeded into six-well plates at a density of 1 million cells per six-well plate, 15 h before transfection. Transfection was performed using Lipofectamine LTX plus (Invitrogen) in Opti-MEM medium. After 6 h of incubation, the medium was changed to advanced DMEM, and cells were cultivated for 24 h at 37°C and 5% CO2. Cell harvesting and RNA isolation were performed using TRI reagent (Sigma-Aldrich).

2.5 Microarray analysis

The entire genome array analysis was performed on an agilent human whole-genome array chip at the Institute of Molecular Biology (University of Mainz). All samples were analyzed in triplets. The R-version 2.13.1 and the extension array QualityMetrics 3.8.0 were used to process the results. Ingenuity and Prism (Graphpad) were the basis of further analyses for expression, networking and survival analysis.

2.6 qPCR Analysis

For qPCR on the targets found by bioinformatics, a two-step strategy was selected. Reverse transcription was performed with the RevertAid H Minus First Strand cDNA Synthesis Kit (Fermentas) using an oligo-dT-primer. Quantitative analysis itself was done with QuantiTect Primer Assays (Quiagen) in a LightCycler LC480 (Roche). Three biological samples were run in triplicate and quantified using a comparative cycle threshold. Further evaluation and $t$-testing were performed with MS Excel 2010.

2.7 Colony-forming assay

Proliferation of the cells was examined with the colony-forming assay. Transfected cells and controls were shown at densities of 5000 and 10000 cells per well. Cells were grown for 1 week, changing media every second day. Colonies were then stained with crystal violet for analysis.

2.8 Impact of CDC3 and KIF18B target gene expression for survival of patients with HCC; predictor development

To investigate the prognostic relevance of CDC3- and KIF18B-dependent target genes in human HCC, we analyzed a dataset containing 53 human genome-wide HCC microarrays (Andersen et al., 2009) using BRB array tools (http://lincs.ncl.gov; BRB-ArrayTools.html; version 3.8.0). All data were subjected to log2 transformation. After normalization using the median over the entire array, all genes with a percentage of missing data >20% were excluded. The remaining genes were filtered for either the CDC3A or the KIF18B target gene list. Unsupervised hierarchical clustering using Euclidean distance and average linkage were used to split the human dataset into two subgroups: A and B (cluster 3; Eisen et al., 1998). For comparing the difference in survival between the two
subgroups, we performed Kaplan–Meier survival analysis and the log-rank test using the MedCalc software packages (http://www.medcalc.be).

To develop a CDCA3- and KIF18B-dependent predictor, we calculated the average expression for each gene in both cluster groups of the training dataset. As the next step, independent test data from 242 patients with HCC (Roessler et al., 2010) were correlated to this predictor by means of Pearson’s correlation. Patients were assigned to one of the two groups (A or B), depending on the higher correlation value. Survival for patients in the test data subgroups was again analyzed by plotting Kaplan–Meier curves (Supplementary Fig. S1).

3 RESULTS

3.1 Bioinformatics co-expression analysis of a large oncogenic microarray dataset, meta-analysis of microarray data and application of a co-regulation approach to large-scale oncogenic data

To identify co-regulated genes and networks, we analyzed expression profiles of 20827 (genome wide) genes over 2158 microarray datasets incorporating 163 diverse tumor entities.
small, but also four larger, subclusters containing (number of genes) of these subclusters, we identified several on a Pearson’s coefficient of 0.8, we identified 428 genes j/C21 CC 0.56. On plotting the size j/C21 of all 2158 datasets.

Using Pearson’s CC of |CC| ≥ 0.8, we identified 428 genes (2.1%) as being highly co-expressed (Fig. 2D). Next, we aimed to identify biologically interacting networks within this still large number of 428 highly co-regulated genes. To estimate an optimal cutoff for separating subnetworks on the basis of the initial unsupervised clustering, we performed C index and KL index analysis and determined the optimal numbers of subclusters. As shown in Figure 2A and B, both algorithms pointed toward good separation of clusters when 74 or 75 subclusters were chosen. For further analysis, we separated the large unsupervised cluster of 428 genes into 75 subclusters, based on a Pearson’s coefficient of |CC| ≥ 0.56. On plotting the size (number of genes) of these subclusters, we identified several small, but also four larger, subclusters containing ≥20 genes (cluster #3, 4, 14 and 16). Cluster #4 was by far the largest subnetwork, containing 61 single genes (Fig. 2D and E).

### 3.2 Identification of functionally related subnetworks by means of biological function enrichment analysis

Analyzing the complete cluster of 428 genes that were found to be highly co-regulated, Ingenuity Pathway Analysis revealed that the most prominent bio-function was ‘Cancer’ (P-value: 1.74E-26–8.7E-03, number of genes: 177), thus validating our approach. In addition, other tumor-related bio-functions such as ‘Cell Death’ (7.49E-10–9E-03, 101) and ‘Cellular Growth and Proliferation’ (5.57E-07–9E-03, 121) were identified. Next, subnetworks with high genetic similarity as demonstrated by unsupervised clustering were analyzed in respect of their signaling pathway and biological function enrichment. Among the largest co-regulated subnetworks was a 61-gene-containing network (#4, CC: 0.59–0.88) that had mainly genes related to cell cycle regulation and cancer development as determined by PubMatrix analysis (Becker et al., 2003). Other co-regulated subnetworks were found to contain enrichment of ribosomal genes (#14, 41 genes, CC: 0.66–0.98); these genes are involved in immunity (#16, 42 genes, CC: 0.56–0.96) and immunological events (Fig. 2D).

### 3.3 Robustness of co-regulated genetic subclusters

Given our hypothesis that the co-regulated networks functionally interact and also for the purpose of biological relevance, these networks should be visible on overall analysis and also stable within individual tissues. For this reason, we compared their stability and behavior in HCC (45 samples, 2.1% of the overall data), breast cancer (353 samples, 16.4%) and colon cancer samples (289 samples, 13.4%). An overlap of 255 of the 428 genes was continuously co-regulated in all three tissues. On average, 78.2% of the genes in the four largest subclusters (#4, #16, #3 and #14) were preserved (at least 57.1%). Thus, high coherence of the cluster function was conserved in several tissues (Supplementary Fig. S2).

### 3.4 Advanced functional analysis of a highly conserved oncogenetic subcluster

Among the different clusters, #4 incorporated genes with high enrichment of carcinogenic and cell cycle-regulating gene ontologies. Given the importance of cell cycle regulation for cancer development, we decided to focus our subsequent analysis on this cluster. To better understand the detailed molecular mechanism by which these genes interfere with cell cycle regulation, we evaluated the enrichment of canonical pathways among these 61 genes. Top-ranked genetic pathways included the mitotic transition and checkpoint kinases (IPA analysis, http://www.ingenuity.com, Fig. 4). Among the different clusters, #4 incorporated genes with high enrichment of carcinogenic and cell cycle-regulating gene ontologies. Given the importance of cell cycle regulation for cancer development, we decided to focus our subsequent analysis on this cluster. To better understand the detailed molecular mechanism by which these genes interfere with cell cycle regulation, we evaluated the enrichment of canonical pathways among these 61 genes. Top-ranked genetic pathways included the mitotic transition and checkpoint kinases (IPA analysis, http://www.ingenuity.com, Fig. 4).

To obtain a broad overview about the known biomedical information concerning these 61 genes, we next performed a text-mining analysis using PubMatrix (Becker et al., 2003). A search based on the terms ‘cancer’, ‘tumor’, ‘liver’, ‘carcinoma’, ‘HCC’ and ‘hepatocellular carcinoma’ revealed that the majority of genes (94%) had already been described with respect to (liver) ‘cancer’. This served as the ‘proof of principle’ for our approach. Two genes had not been previously described with respect to carcinogenesis: CDCA3 and KIF18B. However, as CDCA3 and KIF18B were part of this tightly co-regulated oncogenic and cell cycle-regulating network, we proposed as...
a guilt-of-association hypothesis that these two genes may be novel oncogenes essentially linked to carcinogenesis.

3.5 Activation of CDCA3 and KIF18B in multiple tumor entities

To evaluate the importance of CDCA3 and KIF18B in carcinogenesis, we next investigated differential gene expression in four different tumor entities, i.e. breast, lung, ovarian and renal cancer (n = 5, each). Liver cancer expression was further evaluated in a publicly available dataset of 60 patients (Shimokawa et al., 2010). In all five tumor entities, a highly significant upregulation of both CDCA3 and KIF18B expression was registered in comparison with normal tissue.

Specifically, CDCA3 showed on average 227-, 159-, 560- and 16-fold overexpression in breast, lung, ovarian and renal cancer, respectively, compared with normal tissue. For KIF18B, the gene showed on average >100-, 22-, >100- and >100-fold overexpression in breast, lung, ovarian and renal cancer, respectively, compared with normal tissue. This was particularly because of low expression in normal tissue (Fig. 3).

Besides, CDCA3 was overexpressed in 59 and KIF18B was overexpressed in 49 of 60 patients in a Japanese liver cancer patient cohort with a median 3.59- and 2.34-fold change, respectively (Fig. 3).

3.6 CDCA3- and KIF18B-dependent downstream signatures identified by microarray analysis further confirm a role in carcinogenesis

To further confirm the role of CDCA3 and KIF18B as driver genes in carcinogenesis, we performed microarray analyses after 24-h overexpression of both genes in HUH7 cells. Compared with empty expression vector controls (PCI), we identified 143 genes being differentially regulated by CDCA3 overexpression and 440 genes by KIF18B overexpression. Functional annotation of CDCA3- and KIF18B-regulated genes showed significant enrichment of cancer-related genes. Most downstream target genes showed consistent enrichment of major oncogenic pathways for both genes (P = 9.2E-06 – 1.01E-02 among CDCA3 target genes, P = 5.64E-05 – 1.09E-02 among KIF18B target genes). In addition, many of the highest ranked associated network functions (CDCA3: antigen presentation, cell-to-cell signaling and interaction, hematological system development and function, cell death, cellular movement and cell cycle; KIF18B: cell death, cellular development, hematological system development, cell-to-cell signaling and interaction, cellular movement and immune cell trafficking) or molecular and cellular functions (CDCA3: cellular growth and proliferation P = 6.75E-06 – 1.01E-02, KIF18B: cell death 2.06E-06 – 1.10E-02, cellular development 1.77E-05 – 1.10E-02 14) further support key tumorigenic roles of these genes and their associated molecular signaling pathways (http://www.ingenuity.com).

Together, the enrichment of established cancer-related signaling pathways and biological functions indicates that
overexpression of CDCA3 and KIF18B is deeply linked to functional changes in carcinogenesis.

3.7 Cell cycle regulation by CDCA3 or KIF18B overexpression, RT-PCR validation

As cell cycle regulation is a major hallmark of genes involved in carcinogenesis, we investigated the impact of CDCA3 or KIF18B on cell cycle regulation in greater detail (Fig. 4).

3.8 Cell cycle regulation

The importance of cyclin D1, CDK4 CDK2 and p21/CDKN1A for G1/S-phase transition is widely accepted (Sherr, 1996). Disruption of these genes was analyzed. All four genes showed significant downregulation after CDCA3 overexpression (cyclin D1: −0.37, P < 0.01; CDK4: −0.5, P = 0.020; CDK-2: −0.84, P < 0.01; p21/CDKN1A: −0.93-fold, P < 0.01). The residual activity of 7% for p21/CDKN1A, clearly shown to act as a tumor suppressor, may be equated with nearly complete inactivation of the gene locus. Even greater suppression was observed after KIF18B overexpression (cyclin D1: −0.53, P = 0.01; CDK4: −0.68, P < 0.01; CDK2: −0.86, P < 0.01; p21/CDKN1A: −0.94, P < 0.01). Altogether, the overexpression of CDCA3 or KIF18B caused considerable disturbance in the gene expression of G1/S cell cycle stage-regulating genes (Fig. 4). We then looked for key regulators of G2/M transition (Bucher et al, 2008). Consistently, on overexpression of CDCA3 the cells responded with a strong increase in activity of both examined cyclins, cyclins B1 (2.38-fold, P < 0.01) and B2 (1.78-fold, P = 0.03), compared with PCI. Conversely, the activity of the corresponding cyclin opponent WEE-1 was reduced −0.40-fold (P < 0.01). Overexpression of KIF18B exerted an even stronger effect on the upregulation of cyclins B1 (1.42-fold, P < 0.01) and B2 (1.54-fold, P < 0.01). Consistently, WEE-1 expression was decreased −0.21-fold (P = 0.11). However, despite the clear trend, the latter data failed to achieve statistical significance (Fig. 5).

3.9 Tumor suppressor and oncogenes

Besides major cell cycle checkpoints, well-established tumor suppressor genes were significantly downregulated after CDCA3 and KIF18B expression. TP53 [Lee and Muller, 2010; −0.34-fold (KIF18B, P = 0.11) to −0.43-fold (CDCA3, P < 0.01)] and the apoptosis-inducing gene TRAF2 (Takeuchi et al, 1996) were significantly downregulated [−0.62-fold (CDCA3, P < 0.01) to −0.82-fold (KIF18B, P < 0.01)].

Further, the oncogenes RAN [(Rensen et al, 2008) (2.04-(CDCA3, P = 0.01) to 2.07-fold KIF18B, P = 0.01) and TRIM37 (0.74-(CDCA3, P < 0.01) to −0.64-fold KIF18B, P < 0.01)] revealed a marked increase in expression on CDCA3 or KIF18B expression. Additionally, SKP-1, an essential component of the Skp, Cullin, F-box containing (SCF) complex involved in the degradation of WEE-1, was significantly overexpressed [1.29- (CDCA3, P < 0.01) to 1.72-fold (KIF18B, P < 0.1)] (Jia et al, 2009).

3.10 Overexpression of CDCA3 and KIF18B leads to impaired proliferative capacity due to the disturbance of cell cycle regulation, which is a major deregulation in cancer

To further characterize the functional consequences associated with overexpression of CDCA3 and KIF18B in vitro, we assessed proliferative behavior by colony-forming assay. A massive increase in proliferation compared with the PCI controls was observed. Strongest growth was observed in those cells with excessive CDCA3 activity (number of colonies). Notably, overexpression of KIF18B not only changed the incidence of colony formation but also resulted in a morphological change: larger colonies were found compared with those after CDCA3 overexpression (Fig. 5A).

Interestingly, the induction of CDCA3 and KIF18B did not impair the apoptosis response as assessed by fluorescence-activated cell sorting (FACS), underlining the predominant role of these genes in cell cycle regulation (Fig. 5B).

3.11 Prognostic relevance of CDCA3 and KIF18B downstream target predictors

As the data detailed above pointed toward the major role of CDCA3 and KIF18B in carcinogenesis in general as well as in multiple tumor entities, we specifically investigated their prognostic potential in various cancers, especially HCC.

The 143 CDCA3- and 440 KIF18B-dependent genes were integrated into our 53 HCC patient database (training data, Andersen et al, 2009). Unsupervised clustering resulted in two distinct subgroups. On Kaplan–Meier analysis both the
CDCA3-dependent and the KIF18B predictors demonstrated statistically significant prognostic relevance in these two patient groups ($P = 0.04$, $P = 0.01$, Fig. 6). To validate these results, we further correlated two independent datasets of 242 and 81 HCC patients to the mean of each gene from the training data by means of Pearson’s correlation (test data, Fig. 7). Kaplan–Meier analysis again showed two prognostic subgroups with significant differences in survival in both datasets for CDCA3 ($P = 0.0001$, $P = 0.009$). The 242 patient datasets also showed two prognostic subgroups with significant differences in survival for the 242 patient cohort ($P = 0.005$) and a trend to differences in survival for the 81 patient cohort ($P = 0.11$). Early survival showed highly significant differences for CDCA3 ($P = 0.0000$, $P = 0.0001$) and KIF18B ($P = 0.004$, $P = 0.0006$) in both cohorts (Fig. 7).

Finally, we tested our CDCA3 and KIF18B networks for association with prognosis in a meta-analysis of several other tumors using the Oncomine database (http://www.oncomine.org). Besides HCC, these genetic networks were found to be of prognostic relevance in diverse tumor entities, including breast and colon cancers (Supplementary Table S3).

### 4 DISCUSSION

#### 4.1 Bioinformatics combination of guilt-by-profiling and GBA strategies

The availability of large microarray collections of tumor tissue without corresponding normal tissue serves as a rich source of information about carcinogenesis in general, which may not be achieved by conventional microarray analysis (http://expo.intgen.org/geo/home.do). To extract the key biological changes and functions of carcinogenesis, GBA strategies may provide valuable options for the analysis of data (Lee et al., 2004; Wolfe et al., 2005). GBA assumes that genes with related functions tend to share biological features, such as expression patterns (Oliver, 2000). Generally, on a lower scale, GBA is a concept used repeatedly in biology on a gene-by-gene basis.

Gene functions are frequently estimated and discovered using the analogy of other functionally well-characterized genes. On a genome-wide scale, several bioinformatics publications reported on efforts to improve the computational aspects of GBA approaches for predicting gene function (Hishigaki et al., 2001; Pena-Castillo et al., 2008; Tsuda et al., 2005; Wolfe et al., 2005). However, the number of biologically proven predictions based on high-throughput approaches is still small, and the promise of GBA as a general unbiased method for filling in unknown gene function has not come to fruition. With respect to cancer biology, only 34 publications are listed in PubMed (Sayers et al., 2012) when searching for ‘guilt by association’ and ‘cancer’. Furthermore, most of these publications addressed only methodical aspects of the procedure.

Applying a combined guilt-by-profiling and GBA bioinformatics strategy, we successfully analyzed the GSE2109 microarray repository providing data on 2158 microarray tumor samples (http://expo.intgen.org/geo/home.do) to extract the key biological changes and functions of carcinogenesis. GBA strategies may provide valuable options for the analysis of data (Lee et al., 2004; Wolfe et al., 2005). GBA assumes that genes with related functions tend to share biological features, such as expression patterns (Oliver, 2000). Generally, on a lower scale, GBA is a concept used repeatedly in biology on a gene-by-gene basis.

By analyzing genes for co-regulation across a large collection of tumor samples instead of looking at the differences between tumor and corresponding normal tissue, we identified highly
stable oncogenetic networks. We confirmed these findings by three different molecular and functional approaches: bioinformatics combination of guilt-by-profiling and GBA strategies, identification and molecular validation of novel oncogenes, identification of their downstream functional pathways and identification of novel signatures of prognostic relevance in HCC.

Overall, our work demonstrated the potential of integrating intelligent bioinformatics analyses and sophisticated molecular analyses as a highly valuable tool to obtain novel insights into tumor biology, diagnostics and eventually therapy (Fig. 1).

Besides, we were able to successfully validate the high coherence of a key regulatory network in carcinogenesis, not only in overall analysis but also in individual tissues such as liver, breast or colon cancer. This was of particular interest because averaging Pearson’s CC over a large number of arrays and diverse tissues may be limited by the drawback of the potentially high variability of gene expression correlation within these diverse tissues. We demonstrated two important aspects: given the stability of our cluster, this network may be regarded as a key regulator in carcinogenesis. Second, combining GBA studies with a subsequent analysis in a search of highly coherent subnetworks may yield much greater success in translating this sophisticated bioinformatics strategy into cancer biology and also enhance our understanding of many other diseases in the future.

### 4.2 CDCA3 and KIF18B, identified by bioinformatics GBA profiling, are key regulators of carcinogenesis by interfering with cell cycle regulation

It has become clear that not all genes differentially expressed in cancer are truly genes driving the neoplastic process (‘driver’ genes). It is therefore essential to distinguish these genes from sole bystander genes (‘passenger’ genes). One of the most effective strategies is to analyze their interference with well-established molecular changes in cancer biology. We therefore functionally characterized our genes identified through bioinformatics guilt-by-profiling. We provide several lines of evidence for the key carcinogenic role of these genes in many cancer entities, particularly with respect to cell cycle regulation.

Loss of cell-cycle checkpoints are a hallmark of human cancer because they result in permanent genomic alterations, such as deregulation of oncogenes and tumor suppressor genes (Laiho et al., 2003). Our data supported the interaction of both genes with cell cycle checkpoint-regulating genes and a severe disturbance of cell cycle regulation on overexpression of either one (CDCA3 or KIF18B), ultimately leading to tumor growth (Figs 5 and 6).

First, our large-scale bioinformatics pathway analysis demonstrated significant enrichment of cell cycle-coordinating genes due to overexpression of CDCA3 or KIF18B in the hepatoma cell line HUH7.

Second, qRT-PCR analysis confirmed the deregulation of multiple checkpoint kinases and central cell cycle regulation genes, thus validating the obtained microarray data. B-type cyclins, B1 and B2, essential components of the cell cycle regulatory machinery and both closely connected to G2/M progression, were found to be severely deregulated after overexpression of either CDCA3 or KIF18B. Deregulation of these genes and G2/M may have serious consequences on the cell, such as the development of cancer. This is particularly because, during this phase, cells may arrest transiently to allow for the repair of cellular damage. G1/S-phase-regulating genes were also significantly deregulated, further disrupting normal cell cycle control and enhancing tumor development. P21 deficiency was repeatedly shown to be closely linked to carcinogenesis (Garcia-Fernandez et al., 2011; Hawkes et al., 2011). CDK2 inhibition (by p21) has also been linked to liver cancer development (Kim et al., 2009). CDK4 complexes with cyclin D1 are involved in cell cycle control. Again, this complex may be inhibited by p21 and such deregulation is found in liver cancer (Rivadeneira et al., 2010). However, these changes may not only occur in liver cancer but also in many other tumors.

Third, these data were independently validated by performing an in vitro functional colony-forming assay, which also showed significantly greater development of colonies after overexpression of either CDCA3 or KIF18B.

Taken together, these data concerning interference with cell cycle regulation clearly provide a functional explanation for the driving role of carcinogenesis in several cancer types.

### 4.3 Prognostic relevance of KIF18B and CDCA3 target gene signatures

Gene expression signatures constitute a powerful achievement in the development of novel diagnostic tools, such as accurate and unbiased identification of prognostic subclasses and new cellular targets in liver cancer. The signatures must be robust to be useful in clinical therapeutic algorithms. However, the majority of the reported signatures were not confirmed in further independent datasets (Marquardt et al., 2012; Teufel et al., 2012). To demonstrate the predictive strength of our CDCA3 and KIF18B downstream predictors, we evaluated two additional independent datasets derived from 242 and 81 patients with HCC, and were able to prove their prognostic role in these independent test data (Figs 6 and 7).

Furthermore, as we had originally identified the two novel oncogenes and downstream signatures using a tumor-entity-independent/superordinate approach, we further validated our dependent signatures in several other cancer tissues, such as breast and colon cancer. Given the vigor of predictive significance in multiple tumors, we believe that this signature is robust in respect of essential regulation for HCC as well as cancer in general.

### 5 CONCLUSION

Bioinformatics integration of oncogenetic microarray meta-analysis, guilt-by-profiling and GBA strategies are a suitable approach for the identification of novel carcinogenic networks and oncogenes. Using this approach, we identified two novel oncogenes (CDCA3 and KIF18B), demonstrated their deregulation in multiple tumors, functionally characterized the novel oncogenes, identified corresponding downstream targets and showed the robust prognostic relevance of these downstream predictors for HCC as well as several other tumors.
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