Selection of Personalized Patient Therapy through the Use of Knowledge-Based Computational Models That Identify Tumor-Driving Signal Transduction Pathways

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Abstract

Increasing knowledge about signal transduction pathways as drivers of cancer growth has elicited the development of "targeted drugs," which inhibit aberrant signaling pathways. They require a companion diagnostic test that identifies the tumor-driving pathway; however, currently available tests like estrogen receptor (ER) protein expression for hormonal treatment of breast cancer do not reliably predict therapy response, at least in part because they do not adequately assess functional pathway activity. We describe a novel approach to predict signaling pathway activity based on knowledge-based Bayesian computational models, which interpret quantitative transcriptome data as the functional output of an active signaling pathway, by using expression levels of transcriptional target genes. Following calibration on only a small number of cell lines or cohorts of patient data, they provide a reliable assessment of signaling pathway activity in tumors of different tissue origin. As proof of principle, models for the canonical Wnt and ER pathways are presented, including initial clinical validation on independent datasets from various cancer types. Cancer Res; 74(11): 2936–45. ©2014 AACR.

Major Findings

As expected, the Wnt pathway was predicted inactive in normal colon samples and active in 97% of tested colon adenomas and carcinomas (known to be Wnt driven) and in all tested medulloblastomas containing an activating β-catenin mutation. Furthermore, in primary liver cancer, Wnt activity was predicted in 56% of samples containing a β-catenin mutation, against 18% without documented mutations. In breast cancer, Wnt activity was predicted in 30% of basal-type breast cancers versus 7% of other subtypes, confirming expectations based on clinical research.

The estrogen receptor (ER) pathway model predicted inactivity in practically all tested samples or exceptions for 39% of ER-positive breast cancer samples. Furthermore, ER pathway activity was associated with increased disease-free survival compared with patients in which the pathway was predicted inactive, even though the model was not trained for this purpose.

Clinical implementation of our models is expected to enable a more informed choice of therapy and improved prediction of targeted therapy response. Furthermore, it may help to focus search for tumor-driving genomic defects in whole-genome sequencing data to only those genes involved in the aberrantly active pathway(s).

Introduction

Knowledge on intracellular signal transduction pathways governing cancer cell behavior and controlling cell division is rapidly increasing. This development has elicited a paradigm shift toward development of a whole new category of "targeted drugs," aiming to target the aberrant signaling pathway, which drives tumor growth in the individual patient with cancer (1). In contrast with conventional chemotherapy, targeted therapy requires a highly personalized approach to treatment choice, in principle based on predicting treatment response before administering the drug or drug combination of choice.

The anticipated increasing availability of targeted drugs (2) stresses the need for reliable companion diagnostics to predict therapy response, for which identification of the tumor-driving signaling pathway and the underlying defect that causes its aberrant activation is of high importance (3). Unfortunately, currently available tests often lack predictive value with respect to targeted therapy response. In general, these tests demonstrate (over-)expression of key proteins in signaling pathways of interest, e.g., estrogen receptor (ER) or HER2 in breast cancer, or identify DNA mutations (e.g., in the PIK3CA gene) or structural changes (like HER2 coding gene amplification) in genes encoding...
for proteins that directly influence signaling pathway activity. However, though associated with response to targeted drugs (hormonal therapy and trastuzumab in example cases ER and HER2, respectively), such tests do not provide conclusive information on the functional activity status of cases ER and HER2, respectively), such tests do not provide conclusive information on the functional activity status of the associated signaling pathways (4).

Around 10 major oncogenic signaling pathways play a role in tumor growth and metastasis: ER and androgen receptor pathways, phosphoinositide 3-kinase (activated by multiple growth factor receptors, like HER2, EGFR), canonical Wnt, Notch, Hedgehog, TGF-β, NF-κB, VEGF, RAS/RAF/MAPK/ERK, and FGF signaling pathways (5); see also Supplementary Major Oncogenic Signaling Pathways. Whole-genome and transcriptome analysis methods, like DNA and RNA sequencing and microarray technologies, are in principle capable of producing all data necessary to extract information on signaling pathway activity from cancer tissue samples, on the premise that adequate software is available to interpret the highly complex data (6). This has proved to be a tremendous challenge, and most efforts are directed toward identification of genotypic changes, instead of using the transcriptome to provide information on the functional phenotype of the cancer cell—which is determined in concert by both genotype and microenvironment of cancer cells (7).

We describe the development and partial clinical validation of a new type of knowledge-based probabilistic computational modeling framework for oncogenic cell signaling pathways, which enables functional assessment of pathway activity in individual tissue samples based on quantitative transcriptome data as input. The models have been built using Affymetrix HG-U133Plus2.0 microarrays, widely available from Gene Expression Omnibus (GEO, www.ncbi.nlm.nih.gov/geo) and ArrayExpress (www.ebi.ac.uk/arrayexpress); an overview of used datasets is provided in Supplementary Table S3.

As our pathway models are a simplification of signaling pathway biology and as biologic measurements are typically noisy, we modeled signaling pathways in a probabilistic manner using Bayesian networks.

Quick Guide to Equations and Assumptions

Major assumptions

The key assumption in the Bayesian pathway model we present is that functional activity of a signaling pathway is determined by activity of its respective transcription complex, while the latter can be inferred from mRNA expression data of its transcriptional target genes.

Furthermore, we assumed that the oncogenic signaling pathway driving tumor growth is not transiently and dynamically activated, but long term or even irreversibly. Hence, the model was developed for interpretation of a static cellular condition, and complex dynamic pathway features were not incorporated.

Finally, to handle the noisy characteristics of mRNA expression data from microarrays, as well as additional unknown biologic factors, we modeled signaling pathways in a probabilistic manner using Bayesian networks.

Equations

The main equations related to our Bayesian models describe probabilistic relations between different types of nodes. Conditional probability tables describing the relation between transcription complex node $TC$ and target gene nodes $TGi$ are given by Equation 1, and between each target gene node $TGi$ and its respective probeset nodes $PSij$ by Equations 2a and 2b.
we opted for a probabilistic approach, meaning that relationships (i) between transcription complex and target genes and (ii) between target genes and their respective probesets are described in probabilistic terms. Furthermore, we assumed that the oncogenic signaling pathway driving tumor growth is not transiently and dynamically activated, but long term or even irreversibly. Hence, the model was developed for interpretation of a static cellular condition, and complex dynamic pathway features were not incorporated.

Once the Bayesian network model has been built and calibrated for a particular signaling pathway (sketched below and further detailed in Supplementary Methods), the model can be used on microarray data of a new tumor sample by entering probeset measurements as observations in the bottom layer, and inferring backwards in the model the activity probability of the transcription complex. Hence, this latter probability is the primary read-out used to indicate pathway activity, which can be translated into odds of the pathway being active by taking the ratio of the probability of being active versus inactive (i.e., odds are given by \( p/(1-p) \) if \( p \) is the predicted probability of being active).

### Target gene selection

For optimal performance, the Bayesian network models should contain (only) direct target genes of the respective pathways. Unfortunately, pathway databases such as KEGG (www.genome.jp/kegg) and Biocarta (www.biocarta.com) are fairly incomplete and inconsistent on this aspect (8). Hence, we manually selected target genes based on extensive scientific evidence for each individual gene being a direct target gene of the respective transcription complex, including promoter region motif analysis, transcription factor–binding experiments, and differential expression analysis. For Wnt, extensive research at the Hubrecht Institute over the past decades (9–12) culminated in a list of 34 "bona fide" target genes listed in Supplementary Table S1. For ER, we extensively investigated available literature, as detailed further in the Supplementary ER Pathway Target Gene Selection, yielding the 27 genes in Supplementary Table S2. These numbers of genes are on the one hand low enough to give specific results, but large enough to get robust models.

### Model calibration

The probabilistic relations in the Bayesian network models need to be made quantitative to allow for quantitative probabilistic reasoning. To improve generalization behavior across tissue types, we manually set parameters describing the probabilistic relationships (i) between transcription complex and target genes using the following table:

<table>
<thead>
<tr>
<th>Parameters</th>
<th>PS(_{ij}) = low</th>
<th>PS(_{ij}) = high</th>
</tr>
</thead>
<tbody>
<tr>
<td>( TGi ) down</td>
<td>( AHi,j ) + 1</td>
<td>( AHi,j ) + 1</td>
</tr>
<tr>
<td>( TGi ) up</td>
<td>( PLi,j + PHi,j ) + 2</td>
<td>( PLi,j + PHi,j ) + 2</td>
</tr>
</tbody>
</table>

In these tables, the variables \( AHi,j \), \( PHi,j \), \( PLi,j \), and \( PHi,j \) indicate the number of calibration samples with an absent (A) or present (P) transcription complex that have a low (L) or high (H) probeset intensity, respectively. Dummy counts have been added to avoid extreme probabilities of 0 and 1. For more details on the model construction, we refer to the Supplementary Bayesian Network Construction.

### Statistical tests

Statistical tests, generation of Kaplan–Meier curves, and other graphics were performed using R (13). Generally, one-sided tests were used because the expected sign of a relation is known.

### Results

**Initial validation of the Wnt pathway model**

Two instances of the Wnt model were created, a first one for initial proof of concept using cell line data for calibration, and a second one using a larger calibration dataset with patient data, with the advantage of better reflecting the variation encountered across patient samples.

The first Wnt model was calibrated on data from 12 samples of a Wnt abrogation experiment on an LS174T colon cancer cell line (GSE18560; ref. 14), of which six have an active Wnt pathway and six an inactivated Wnt pathway. For initial proof of concept, this model was tested on a dataset with 32 normal colon tissue samples and 32 colon adenoma samples from patients (GSE8671; ref. 11). The Wnt pathway is thought to be active in colon adenoma and inactive in normal intestinal tissue (15), and Fig. 2A shows that this is almost perfectly predicted by our model. Although two of the 32 adenoma samples are predicted to have an inactive Wnt pathway, if the threshold is set at odds of 0.1, the difference with normal colon samples is highly distinctive. Note that reported odds get as large as a million to one, which is due to the model using 34 genes (83 probesets in total), so although one gene may give quite noisy information, combining 34 genes gives quite confident predictions.
Wnt-active surgical colon cancer samples may be explained by cancer tissue heterogeneity and abnormal gene promoter methylation associated with more advanced cancer, or the surgery-associated sampling procedure may have resulted in mRNA degradation and unreliable microarray results (7, 17). Such factors may interfere with the expected Wnt target gene mRNA profile and reduce sensitivity of our current model. Indeed and illustratively, when our model was applied to the transcriptome of Wnt-active colon cancer cell line HCT116, in which specific Wnt target genes are methylated (18), the model predicted an inactive Wnt pathway (odds 8:1, data not shown).

Above results provide evidence that the model can identify active versus inactive Wnt pathway state in tumors arising from the colon epithelial cell type used to develop and calibrate the model.

**Use of the Wnt model in tumors of other tissue origin**

The aim of our models is to enable wide diagnostic usage across tumors of different cellular origins. Although direct target genes are transcribed by induced activity of one or more pathway-specific transcription factors binding to their respective gene response elements, indirect target genes are more likely to depend on additional cellular proteins for their transcription, increasing the likelihood of cell type–specific effects on expression regulation. For this reason, gene selection used to build the models focused on direct pathway target genes. To evaluate this premise of relative tissue-type independent functioning, we subsequently tested the model on other tumor types.

**Liver cancer**

In hepatocellular carcinoma (HCC) and hepatoblastoma, heterozygous somatic mutations (or deletions) in the third coding exon (codons 32, 33, 34, 37, 41, and 45) of the β-catenin (CTNNB1) gene have been frequently identified, resulting in substitution (or loss) of an amino acid, which may be associated with aberrant activation of the Wnt pathway (19–21).

Dataset GSE9843 (22) contains 91 liver cancer tissue samples, of which 27 contain a β-catenin gene mutation, 60 possess wild-type β-catenin, and four are unknown. Furthermore, 31 of the 91 samples scored positive for nuclear β-catenin staining, 55 negative, and five unknown. Interestingly, correlation between β-catenin mutation status and nuclear staining is not significant in this dataset (OR = 2.3; one-sided Fisher exact test, \( P = 0.07 \)), illustrating the difficulty to get ground truth information on Wnt pathway activity in these samples, due to lack of a reliable test.

Chiang and colleagues (22) applied hierarchical clustering based on mRNA microarray data of the 91 samples of GSE9843, yielding five groups labeled “unannotated,” “polysomy chr7,” “inflammation,” “proliferation,” and “CTNNB1.” Despite its label, the latter group of 24 samples only contains 16 of the 27 samples with a β-catenin mutation, and 14 of the 31 samples with a positive staining. The results of our Wnt pathway model on this dataset are shown in Fig. 3A. In the CTNNB1 group, 83% (20 of 24) of the samples are predicted to have an active Wnt pathway, versus 26% (6 of 23) in the proliferation group and 0% in the other three groups. In addition, although β-catenin mutation status and staining were not significantly correlated,
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we found a significant correlation of Wnt pathway activity with both β-catenin mutation status and β-catenin staining (OR = 5.4 and 6.1; respectively; and one-sided Fisher exact test, \( P = 6.9 \times 10^{-5} \) and \( 3.4 \times 10^{-4} \), respectively; detail in Supplementary Detailed Liver Cancer Results on GSE6843).

Analysis of data from a second dataset of hepatocellular carcinoma samples (GSE6764; ref. 23) with unknown mutational status yielded an increased incidence of Wnt activity in patients with relatively more malignant tumors, as seven of the 27 early HCC, advanced HCC, and very advanced HCC samples were predicted to have an active Wnt pathway, compared with none in the very early HCC and nonmalignant sample groups (Fig. 3B, one-sided Fisher exact test, \( P = 4.5 \times 10^{-7} \)).

Medulloblastoma

In medulloblastoma, a subset of tumors is known to possess an activating mutation in the β-catenin gene (24). We applied our model in a blinded manner to the medulloblastoma dataset from Kool and colleagues (\( n = 62 \), GSE10327; ref. 24), and successfully identified all samples with a Wnt pathway-activating β-catenin mutation (Fig. 4). In another medulloblastoma dataset (\( n = 40 \), GSE12992; ref. 25), the Wnt model also correctly identified the four samples with a driving β-catenin mutation against 36 without (data not shown). These two datasets show perfect performance of the model in this tumor type with 100% sensitivity and 100% specificity.

Breast cancer

In patients with breast cancer, direct Wnt-activating gene mutations are generally not present, except for a few rare metaplastic breast cancer cases (26). Rather, activation of the Wnt pathway has been indicated by circumstantial evidence in patients with triple-negative or basal-type breast cancer, most likely induced by interaction between the cancer cell and its microenvironment (27).

Two datasets were used for analysis of Wnt pathway activity within breast cancer subtypes: GSE12276 (28) and GSE21653 (29); see Fig. 5A and B. Using mRNA data, patient samples in these datasets were subtyped according to Desmedt and colleagues (30) and Perou and colleagues (31), respectively. Despite the difference in subtyping approach, Wnt pathway activity was very comparable, with an active Wnt pathway in 30% (21 of 65 and 21 of 75, respectively) of basal-type cancer samples, versus only 7% (15 of 139 and 9 of 191, respectively) in other breast cancer subtype samples (one-sided Fisher exact test, \( P = 2.7 \times 10^{-4} \) and \( 4.8 \times 10^{-7} \), respectively).

Taken together, the above results provide initial evidence that the Wnt model performs well on a variety of tumors without requiring additional training steps on the different cell types of origin.

Initial validation of the ER pathway model

The ER pathway model was calibrated on data from eight samples of the breast cancer cell line MCF7, of which four were deprived from estrogen and four were stimulated with 25 nmol/L E2 (GSE8597; ref. 32), yielding an inactive and an active ER pathway, respectively. Estradiol concentrations are typically around 0.5 nmol/L in normal breast tissue, but elevated in breast cancer to around 2 nmol/L (33). As a result, the 25...
The ER pathway model analysis on mRNA microarray data from samples of patients with breast cancer from datasets GSE12276 (E, blue) and GSE21653 (F, ref. 29) subtyped according to Perou’s subtyping scheme (31), as luminal A (green), luminal B (dark blue), HER2 (orange), and basal (red). All patients within this dataset suffered a relapse (median time to recurrence, 21 months; range, 0–115 months). The ordering of samples within each group is different in the two graphs; only four of the 266 samples have both an active Wnt and ER pathway. B and D, odds of Wnt (B) and ER (D) pathway activity in a dataset with breast cancer samples (Fig. 5E and F), we observed in datasets GSE12276 (E, n = 204; ref. 28) and GSE21653 (F, n = 266; ref. 29). Samples are grouped here according to ER IHC status as ER negative (red), ER positive (blue), or unknown (orange).

Next, we applied the ER pathway model to the same two datasets of patients with breast cancer used above: GSE12276 (28) and GSE21653 (29). Fig. 5C and D, respectively. The figures show an active ER pathway in 41% (38 of 92) and 61 of 138, respectively) of luminal-type patients, versus only 4% (3 of 102 and 7 of 128, respectively) in other breast cancer subtype samples (one-sided Fisher exact test, \( P = 1.3 \times 10^{-10} \) and \( 3.7 \times 10^{-14} \), respectively). The five HER2-type patients with predicted active ER pathway had scored positive for ER and/or progesterone receptor (PR) by immunohistochemistry (IHC) staining. All basal-type breast cancer samples were predicted to have an inactive ER pathway.

If patient samples are grouped according to ER IHC status (Fig. 5E and F), we observed in datasets GSE12276 and GSE21653 an active ER pathway in 39% (27 of 88 and 67 of 150, respectively) of ER-positive tumors, versus practically none (1 of 77 and 0 of 113, respectively) of the ER-negative tumors.

Figure 5. Results from Wnt (A and B) and ER (C–F) pathway model analysis on mRNA microarray data from samples of patients with breast cancer. The y-axis shows the odds of the respective pathway being active (“on”) versus inactive (“off”) on a logarithmic scale. Each tissue sample result is represented by a bar. A and C, odds of Wnt (A) and ER (C) pathway activity in a dataset with breast cancer samples (\( n = 204 \), GSE12276; ref. 28) subtyped according to the module approach from Desmedt and colleagues (30) as luminal A (green), luminal B (dark blue), HER2 (orange), and basal (red). All patients within this dataset suffered a relapse (median time to recurrence, 21 months; range, 0–115 months). The ordering of samples within each group is different in the two graphs; only four of the 204 samples have both an active Wnt and ER pathway. B and D, odds of Wnt (B) and ER (D) pathway activity in a dataset with breast cancer samples (\( n = 266 \), GSE21653; ref. 29) subtyped according to Perou’s subtyping scheme (31), as luminal A (green), luminal B (dark blue), HER2 (orange), and basal (red). All patients within this dataset suffered a relapse (median time to recurrence, 21 months; range, 0–115 months). The ordering of samples within each group is different in the two graphs; only four of the 266 samples have both an active Wnt and ER pathway. E and F, results from the ER pathway model analysis on mRNA microarray data from samples of patients with breast cancer from datasets GSE12276 (E, n = 204; ref. 28) and GSE21653 (F, n = 266; ref. 29). Samples are grouped here according to ER IHC status as ER negative (red), ER positive (blue), or unknown (orange).
ER pathway activity and tamoxifen sensitivity

To link ER pathway activity to tamoxifen sensitivity, we analyzed dataset GSE21618 (37), containing samples from tamoxifen-sensitive and -resistant MCF7 breast cancer cell lines (all ER positive) treated with estradiol. In particular, we took the samples that had first been deprived of estrogen, and next been stimulated for up to 48 hours. Figure 6 shows the resulting probability of ER pathway activity as a function of stimulation time. Clearly, tamoxifen-sensitive cell lines quickly respond to estrogen stimulation, with probabilities steeply increasing toward 1, whereas tamoxifen-resistant cell lines respond to a lesser extent.

Initial assessment of prognostic value of the ER pathway model

Although the pathway models have been developed and trained to assess pathway activity to predict therapy response, we also tested to what extent they can have prognostic value. To this end, survival time analysis was performed on a dataset of 164 patients with ER-positive breast cancer that all received (only) adjuvant tamoxifen treatment for 5 years (Fig. 7; combined datasets GSE6532 & GSE9195; refs. 38, 39). The analysis was restricted to the first 5 years only, as tamoxifen treatment was limited to 5 years. As expected, patients with an active ER pathway have a better survival prognosis on tamoxifen treatment than patients for which the ER pathway is predicted inactive (one-sided log-rank test $P = 0.034$).

These results indicate that the ER pathway model may also have clinical utility in assessing prognosis in individual patients with breast cancer, even though it has not been developed for this purpose.

Discussion

With a few exceptions, e.g., HER2 and ER protein staining in breast cancer, most companion diagnostic assays to predict therapy response focus on identification of a tumor-specific genetic defect, associated with activation of a specific oncogenic signaling pathway (4, 40, 41). If the test result is positive, the associated signaling pathway is assumed to be active and driving tumor growth. Although a recent study by MD Anderson provides proof of principle that mutation-based identification of the tumor-driving signaling pathway improves therapy choice, it also illustrates that such a DNA-based companion diagnostic approach is unlikely to provide the complete answer, as in this study therapy response increased from 5% to only 27% (41). Indeed, it is clear that signaling pathway activation status (the functional “phenotype” of the cell) is determined not only by errors in the cancer cell (epi-)genome, but to a large extent by interactions between the cancer cell and its microenvironment (1, 32, 42). To assess the phenotype of the cell, in addition to the genotypic, we presented a method to interpret cancer tissue transcriptome data as direct quantitative “output” of active signal transduction pathway(s), using knowledge-based Bayesian models. The pathway “output” is represented by transcribed pathway-specific target genes, which need to be known to create the models. So far, the data input for the models is from Affymetrix HG-U133Plus2.0 microarrays, but an important advantage of this type of knowledge-based models is that they can be easily calibrated to other input modalities, such as other array types, RNA sequencing, or dedicated multiplex PCR assays. We provided proof of principle that the models when trained on a limited number of cell line samples, or if available patient samples, already perform very well and robustly identify active oncogenic signal transduction pathway(s) in individual tissue samples obtained from a variety of malignancies.

For the Wnt pathway, our pathway model analysis results for adenoma, colon carcinoma, and medulloblastoma were in full concordance with existing evidence on Wnt pathway activation in these tumor types (15, 24, 42).

For other tumors such as primary liver and breast cancer, no easy “ground truth” with respect to activity status of the Wnt
pathway is available, but this pathway is likely to play a role in at least a number of cases (42–45). In agreement, our model identified an active Wnt pathway in the majority of liver cancer samples with, and a minority without, a β-catenin mutation. Identification of an active Wnt pathway in the absence of a β-catenin mutation may be due to the presence of other pathway-activating mutations in genes like APC, AXIN1, and AXIN2 (46), or by paracrine Wnt activation (47, 48). Indeed, a significant correlation was found between model-predicted Wnt pathway activity and staining of nuclear β-catenin, presumably the active form of the Wnt pathway transcription factor. On the other hand, the presence of a β-catenin mutation does not necessarily mean that the pathway is activated, and in the cases in which the model did not detect Wnt activation despite a β-catenin mutation, another pathway may have taken the lead in tumor growth, for example induced by the microenvironment.

In breast cancer, Wnt activity was detected by the model in around one third of triple-negative or basal-type samples, which agrees with available evidence on a role for Wnt activity in this breast cancer subtype (43). In only very rare cases of breast cancer, a potentially Wnt-activating gene mutation has been found, suggesting that Wnt activity is most likely induced by paracrine interactions between cancer cells and their microenvironment. With a number of Wnt-targeting drugs in the pipeline of pharmaceutical companies, development of a reliable test to identify Wnt pathway activity in this cancer subtype with highly unfavorable prognosis is considered high priority (44, 49, 50), as β-catenin staining is not reliable enough to indicate Wnt pathway activity (50). Analysis of transcriptome data by our model is expected to provide information on Wnt activity in breast cancer, but final validation will require a clinical trial with an appropriate Wnt-inhibiting drug.

Other approaches have been directed toward assessing the phenotype of cancer by analyzing its transcriptome. To deduce pathway activity from tissue transcriptome data, most pathway analysis approaches use pathway information from databases such as KEGG (www.genome.jp/kegg) and Biocarta (www.biocarta.com) that mainly list genes encoding signaling proteins. Furthermore, they invariably extrapolate quantitative mRNA levels to levels of corresponding signaling proteins, followed by a search for a role of the transcript-encoded protein in a signaling pathway, which is subsequently defined as an active pathway (42, 44, 51). This approach is intrinsically flawed for pathway activation analysis because induction of an mRNA transcript coding for a component of a signaling pathway is not reliably correlated to the actual translated protein level and, even less, to the activation status of the encoded signaling protein, which requires additional post-translational protein modifications.

As expected, our approach leads to a different interpretation result of mRNA profiling data than conventional pathway analysis approaches. For example, we have run gene set enrichment analysis (52), using the accompanying curated canonical pathways, to identify the pathways differentially activated between the normal colon and colon adenoma samples from dataset GSE8671 (11), but none of the Wnt-related pathways was identified with a significant P value after correction for multiple testing (see Supplementary Table S5 in Supplementary Methods). Furthermore, another common pathway analysis approach, as presented by Skrzypczak and colleagues (16), in which first a list of differentially expressed genes is determined and next pathway sets are analyzed for overrepresentation of this gene list, did not identify the Wnt pathway as significantly different between normal colon and colon neoplasms—while our results were highly convincing about Wnt pathway activation status.

From the results on tamoxifen-sensitive and -resistant cell line data, it is inferred that the ER model can successfully detect ER activity in breast cancer cells. Being positive for the ER as measured by IHC or microarray analysis seems to be a necessary but not sufficient condition for ER pathway activity as assessed by the model. Because the model was trained on data from cell line experiments performed with a relatively high dose of estradiol (25 nmol/L), it cannot be excluded that the current model lacks to some extent in sensitivity to detect all samples with an active ER pathway. However, the finding that only a subgroup of ER-positive patients seemed to have an active ER pathway agrees with the common clinical observation that a number of ER-positive patients are primary resistant to hormonal therapy. Moreover, the model identified ER-positive patients treated with hormonal adjuvant therapy but with an inactive ER pathway as having a worse prognosis. This is in agreement with the concept that patients with the ER pathway driving tumor growth are more likely to benefit from hormonal adjuvant therapy.

With respect to the prediction of hormonal therapy response in patients with breast cancer, Symmans and colleagues (53) have described an mRNA profile of 165 ER-related genes, called the sensitive to endocrine therapy (SET) index. In contrast with the ER target genes in our model, genes underlying the SET index were not selected based on evidence for them being target genes of the ER transcription factor, but on increased expression levels found in ER-positive patients. The SET index was shown to identify patients with node-negative disease that have a good prognosis when treated with adjuvant hormonal therapy, as well as patients treated with neoadjuvant chemotherapy who have a high risk at relapse when subsequently treated with adjuvant hormonal treatment. Its predictive value may be partly contributed to the incorporation of six of the ER target genes used in our model.

We hypothesize that in ER-positive patients with an inactive ER pathway, other pathways such as Wnt or Hedgehog, associated with more aggressive behavior and worse outcome, may actually have been driving tumor growth (1, 54, 55). This is in agreement with the reported decline in SET index with advancing pathologic cancer stage, despite ER positivity, suggesting decreasing tumor dependency on an active ER pathway (53). Second, a high gene expression grade index, developed in a similar way as the SET index, also identifies subpopulations of ER-positive breast cancer with unfavorable prognosis (56). Taken together, these results strongly suggest that conventional quantification of nuclear ER IHC staining is not sufficiently specific in detecting functional ER pathway activity. According to pathologist guidelines, ER activity in a breast cancer sample is inferred from the presence of positive ER staining, with a minimum of 1% of ER-positive tumor nuclei as a threshold level. Such staining assays have an estimated 20% error rate due to
multiple factors that influence reliability, among them nonstandardized staining procedures and subjective interpretation (42, 50, 57). In contrast, microarray assays are quite standardized, and the model-based analysis uses a number of ER target genes interpreted in a weighted manner to calculate a probability of pathway activation, instead of ER as single variable as is the case in IHC testing. These conditions are promising for a more robust prediction of ER pathway activity.

Clinical utility of pathway models

The fact that the Bayesian network models are knowledge- and not data mining-based has several advantages. First, the models seemed to be well applicable to data analysis of multiple unrelated tumor types and this property allows use of the models for diagnostics purpose in cancers of different cell types of origin. Nevertheless, it is expected that the models can be further optimized with respect to sensitivity and specificity by including target genes that are specifically expressed in the tissue type of origin of the tumor. Such adaptation could, for example, entail adjusting the conditional probabilities and/or adding new nodes for novel target genes. Another advantage is the relatively easy translation of the model to another data format, such as Illumina DASL microarrays, RNA sequencing, or PCR-based testing. Finally, the remarkable reproducibility of the pathway model results across multiple (mostly public) datasets of a specific tumor type, generated at completely different hospitals and locations, demonstrates the robustness of this modeling approach for individual patient diagnostic use. The pathway model series will be extended to include all major oncogenic signaling pathways, ultimately providing a multipathway analysis suite for identification of both the major active pathway and potential underlying resistance pathways, applicable to tissue samples from a number of tumor types. Upon further clinical validation, currently under way, the expected main clinical utility of the described pathway models lies in therapy response prediction and monitoring in neoadjuvant and metastatic settings.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

Authors’ Contributions

Conception and design: W. Verhaegh, A. van de Stolpe
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Acquisition of data (provided animals, acquired and managed patients, provided facilities, etc.): R. Versteeg, J. Martens, J. Foekens, H. Clevers
Analysis and interpretation of data (e.g., statistical analysis, biostatistics, computational analysis): W. Verhaegh, H. van Ooijen, M.A. Inda, M. Smid
Writing, review, and/or revision of the manuscript (i.e., reporting or organizing data, constructing databases): R. Versteeg, J. Foekens
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