Genome-wide Analysis of Therapeutic Response Uncovers Molecular Pathways

Governing Tamoxifen Resistance in ER+ Breast Cancer

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ABSTRACT

Despite recent advances in diagnosis, classification, and therapeutic management, breast cancer (BC) remains one of the leading causes of cancer-related death in women worldwide. Nearly 70% of all diagnosed cases of breast tumors are Estrogen Receptor positive (ER+) and thus anti-estrogen therapy, such as tamoxifen, has become the standard-of-care for patients with ER+ breast cancers. Yet, nearly 30% of patients treated with tamoxifen develop resistance, ultimately leading to metastasis and lethality. Prioritization of breast cancer patients based on the risk of resistance to tamoxifen plays a significant role in personalized therapeutic planning and improving disease course and outcomes. In this work, we demonstrate that a genome-wide pathway-centric computational framework elucidates molecular pathways as markers of tamoxifen resistance in ER+ breast cancer patients. Through the association of pathway activity and response to tamoxifen, we identified five biological pathways and demonstrated their ability to predict the risk of tamoxifen resistance in two independent patient cohorts (Test cohort1: log-rank p-value = 0.02, adjusted HR = 3.11; Test cohort2: log-rank p-value = 0.01, adjusted HR = 4.24). Importantly, as a negative control, we have demonstrated that the identified 5 candidate pathways did not classify patients simply based on the disease aggressiveness and that pathways of aggressiveness do not overlap with the 5 candidate pathways. Finally, we have compared our pathway signature to other known signatures of tamoxifen response and have shown superiority of our pathway-based approach (adjusted hazard ratio = 3.11, hazard p-value=0.0278). Thus, we propose that the identified pathways as well as their representative read-out-genes can be utilized to prioritize patients who would benefit from
tamoxifen treatment and patients at risk of tamoxifen resistance that should be offered alternative regimens.
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Chapter I

INTRODUCTION

1.1 Background, Literature Review and Statement of the Problem

Despite recent advances in diagnosis, classification, and therapeutic management, breast cancer (BC) remains one of the leading causes of cancer-related death in women worldwide. The main molecular subtypes of BC are based on the gene expression profiling, including luminal A, luminal B, HER2-enriched, and triple-negative/basal-like (Table 1). The luminal type of breast cancer, which is mainly categorized by estrogen receptor-positive (ER+), is morphologically well-differentiated and displays a comparatively good prognosis compared with ER− breast cancers which tend to be poorly differentiated and display a poor prognosis. However, the studies have shown that 65–70% of breast cancers are luminal A and B tumors, whereas about 10% of breast cancers are HER2-enriched tumors and 10–19% of breast cancers are basal-like tumors. Those molecular classifications of BC subtypes significantly enhanced our understanding of the complicated properties of various breast tumors, their clinical overall outcomes, and their treatment responses.
Table 1.
Different Characteristics of breast Cancer subtypes.

| Breast Cancer Molecular Subtypes                  | Characteristics                                      |
|--------------------------------------------------|------------------------------------------------------|
| Luminal A                                        | ER+ and/or PR+, HER2- and Ki-67 ≤14%                 |
| Luminal B                                        | ER+ and/or PR+, HER2+-/any Ki-67 >14%/any Ki-67       |
| HER2-Enriched/Non-Luminal                        | ER-, PR-, HER2+ and any Ki-67                        |
| Triple Negative/Basal-Like                       | ER-, PR-, HER2- and any Ki-67                        |

Nearly 70% of all diagnosed cases of breast tumors are ER+, making treatments with anti-estrogen effects in the breast cells, such as tamoxifen, the standard-of-care for patients with ER+ breast cancers. Despite the significant success of tamoxifen administration, nearly 30% of treated patients develop therapeutic resistance, ultimately leading to metastasis and lethality. Therefore, prioritization of patients based on the risk of resistance to tamoxifen before treatment administration could play a significant role in personalize therapeutic planning for patients with ER+ breast cancer and builds a foundation to improve disease course and outcomes. In addition, identifying which patients are not likely to develop tamoxifen resistance is equally as important as identifying which patients are at high risk for developing tamoxifen resistance.
Tamoxifen is a selective estrogen receptor modulator (SERM) and has agonist or antagonist activity depending on the tissue type. In the breast cells, tamoxifen directly binds to the ER, blocking estrogen from attaching to the receptor and thus inhibiting the activity of estrogen-regulated genes and causing the repression of estrogenic effects. However, the emergence of alternative mechanisms of estrogenic stimulation has been shown to cause emergence of resistance to tamoxifen. For example, some studies have demonstrated that ER+ breast cancers that overexpress HER2 and EGFR can activate the components of downstream signaling pathways which then stimulate both ER and estrogen receptor co-activator AIB1, and thus induce the estrogen agonistic activity of tamoxifen in breast cancer cells. Another study noticed that the increased expression of HER2 signaling can also downregulate progesterone receptor (PR) levels in the ER+ breast tumors, where losing the PR expression serves as a biomarker of hyperactive growth factor signaling, leading to another possible mechanism of tamoxifen resistance. Despite the emerging role of HER2 in tamoxifen resistance, it only accounts for 10% of ER+ breast cancers, suggesting more complex resistance mechanisms in these cases, presenting a central clinical problem for patients with ER+ breast cancer.

In recent years, several groups have developed gene expression signatures of tamoxifen response for ER+ patients, including 10 gene-signature by Men et al., 21 gene-signature by Paik et al. (known as Oncotype DX) and 2 gene-signature by Ma et al. While these signatures provide substantial advances to our understanding of individual genes involved in resistance, they do not yet capture the complex interplay between biological mechanisms that governs tamoxifen resistance. Here we propose a pathway-centric computational framework to elucidate tamoxifen resistance and demonstrate that it
outperforms known gene-based approaches. Advantages of our pathway-based approach lies in (i) its ability to identify a tightly connected cooperative group of genes unified by the same function;\textsuperscript{21-23} (ii) studying molecular pathways, rather than individual genes, produces more reliable read-out outputs as they are less susceptible to experimental noise;\textsuperscript{24} (iii) pathway-level view enhances our understanding of the biological mechanisms related to disease and treatment response;\textsuperscript{25-28} and finally (iv) looking at alterations in biological pathways enhances the likelihood of identifying potential therapeutic targets to preclude or overcome resistance.

1.2 Research Hypotheses

We suggest that the defined candidate pathways as well as their representative read-out-genes can potentially be used to identify patients who would benefit from the tamoxifen treatment as their first-line therapy and those at risk of developing therapy failure, even prior to treatment administration, which enhances the personalized and precision treatment strategy. In fact, identifying which patients are not likely to develop tamoxifen resistance is equally as important as identifying which patients are at high risk for developing tamoxifen resistance. While the identified molecular pathways act as promising predictive markers for treatment response, they can also be potential candidates for therapeutic target to prevent resistance. Although this work is focused on identifying patients with high potential to antiestrogen resistance, our approach can be broadly applicable to other therapeutic interventions and diseases.

1.3 An Overview of the Study
In this work, we have established a systematic pathway-centric computational framework to elucidate molecular pathways as markers of tamoxifen resistance in ER+ breast cancer patients. Through the analysis of pathway activity in each ER+ patient and their association with response to tamoxifen \((n = 53)\), we identified five biological pathways as pathways essential for tamoxifen resistance: Retrograde Neurotrophin Signalling, Loss of NLP from Mitotic Centrosomes, RNA Polymerase III Transcription Initiation from Type 2 Promoter, EIF2 pathway, and Valine, Leucine and Isoleucine Biosynthesis. We have demonstrated the ability of the identified five \((5)\) candidate pathways to predict the risk of tamoxifen resistance in two independent patient cohorts\(^{29}\) (Test cohort 1, \(n = 66\) : log-rank p-value = 0.02, accuracy in leave one out cross-validation \((\text{LOOCV}) = 85.8\%\); Test cohort 2, \(n = 77\): log-rank p-value = 0.01, accuracy in LOOCV = 82.5\%) and their independence from known covariates, such as age, tumor grade, tumor size, lymph node status, and PR status, as the absence of PR in ER+ tumor can be an indicator of HER2 activation and an aggressive phenotype\(^{16}\) (Test cohort 1, adjusted hazard ratio = 3.11; Test cohort 2, adjusted hazard ratio = 4.24). In addition, we have shown significant non-random predictive ability of our pathways, when compared to pathways chosen at random \((\text{random model } p\text{-value}=0.031)\). Furthermore, we performed stratified Kaplan-Meier survival analysis, where we evaluated predictive ability of our candidates on patient groups divided by PR status, age groups, and luminal subtypes and demonstrated that the five candidate pathways can predict risk of resistance to tamoxifen in each group. Importantly, as a negative control, we have demonstrated that the identified five candidate pathways did not classify patients simply based on the disease aggressiveness \((\text{log-rank } p\text{-value} = 0.7, \text{hazard ratio} = 1.246)\) and that in fact pathways associated with disease
aggressiveness do not overlap with the five candidate pathways. We have compared our method to other computational techniques to tackle treatment response, including Epsi et al.\textsuperscript{27} (which utilized extreme-responder analysis, using tails of the treatment response distribution to define a treatment response signature), Zhong et al.\textsuperscript{30} (which used Support Vector Machine approach as a base), and Yu et al.\textsuperscript{31} (which uses random forest approach as a base) and demonstrated that our method outperforms these techniques in predicting risk of resistance to tamoxifen. Further, we have compared our pathway signature to other known signatures of tamoxifen response\textsuperscript{18-20} and have shown the superiority of our pathway-based approach (adjusted hazard ratio = 3.11, hazard p-value = 0.0278). Finally, we identified 5 read-out genes that can function as biomarker to identify patients at risk of developing resistance to tamoxifen in testing cohort. Thus, we propose that the identified five candidate pathways and their corresponding read-out genes can potentially be used to prioritize patients who would benefit from tamoxifen treatment as their first-line therapy, and to identify patients at risk of tamoxifen resistance who should be offered an alternative regimen plan.
Chapter II

METHODS

2.1 Patient cohorts utilized in this study

All gene expression datasets of patients with ER+ breast cancer were obtained from publicly available GEO data repository\textsuperscript{32} from multi-institutional multi-PI comprehensive Loi et al.\textsuperscript{29} study GSE6532 (Figure 1, Table 2): (i) KIT-GSE6532 utilized as a Training cohort; (ii) GUYT-GSE6532, utilized as Test cohort 1; (iii) OXFT-GSE6532, utilized as a Test cohort 2; and (iv) KIU-GSE6532, utilized as a negative control cohort. Training cohort contains patient profiles of primary ER+ breast tumors \((n = 57)\), archived at the Uppsala University Hospital (Uppsala, Sweden), profiled on Affymetrix Human Genome U133A array and Affymetrix Human Genome U133B array. Test cohort 1 contains patient profiles of primary tumors from patients with ER+ breast cancer \((n = 70)\), archived at the Guy’s Hospital (London, United Kingdom), profiled on Affymetrix Human Genome U133 Plus 2.0 Array. Test cohort 2 contains patient profiles of primary ER+ breast tumors \((n = 77)\), archived at the John Radcliffe Hospital (Oxford, United Kingdom), profiled on Affymetrix Human Genome U133A, B array. Negative control cohort consists of not-treated patients with ER+ primary breast tumors \((n = 51)\), profiled on Affymetrix Human Genome U133A, B array. All primary tumors samples in Training and Test cohorts collected through surgery, diagnosed between 1980 and 1995 and received tamoxifen-only treatment for 5 years post-diagnosis as their adjuvant treatment.
Figure 1.

Schematic representation of the utilization of independent patient cohorts for Training, Testing, and negative-control purposes utilized in this study.
Table 2.

Clinical characteristics of datasets used for Training and Testing analysis

| Characteristics | Training cohort KIT-GSE6532 | Test cohort 1 GUYT-GSE6532 | Test cohort 2 OXFT-GSE6532 | Negative-control cohort KIU-GSE6532 |
|-----------------|-----------------------------|----------------------------|----------------------------|-------------------------------|
| Platform        | Affymetrix Human Genome U133A, B array | Affymetrix Human Genome U133 Plus 2.0 Array | Affymetrix Human Genome U133A, B array | Affymetrix Human Genome U133A, B array |
| Total number of patients | 57 | 70 | 77 | 51 |
| PAM50 Classification | | | | |
| Luminal A | 45 | 27 | 61 | 43 |
| Luminal B | 8 | 39 | 16 | 4 |
| *Other subtypes | 4 | 4 | - | 4 |
| Number of patients utilized in our study (other subtypes excluded) | 53 | 66 | 77 | 47 |
| Number of events | 22/53 (41.5%) | 19/66 (28.78%) | 20/77 (25.97%) | 17/47 (36.17%) |
| Age | | | | |
| =<50 | 1 | 3 | 13 | 19 |
| >50 | 52 | 63 | 64 | 28 |
| Histological grade | | | | |
| 1 | 10 | 17 | 17 | 22 |
| 2 | 37 | 36 | 46 | 20 |
| 3 | 6 | 13 | 14 | 5 |
| Tumor size       | 19 | 35 | 35 | 26 |
|------------------|----|----|----|----|
| ≤2 cm            |    |    |    |    |
| >2 cm            | 34 | 31 | 42 | 21 |

| Lymph node status | 14 | 22 | 48 | 47 |
|-------------------|----|----|----|----|
| Negative          |    |    |    |    |
| Positive          | 39 | 44 | 29 | -  |

| PR status         | 4  | 16 | -  | 1  |
|-------------------|----|----|----|----|
| Negative          |    |    |    |    |
| Positive          | 49 | 50 | -  | 46 |

Note: * other subtypes = HER2-enriched, basal-like, or normal-like.

2.2 Data normalization

For each gene expression microarray dataset, matrix of RMA (Robust Microarray Analysis) normalized signal intensity values were used. Using the current annotation file from GEO and the latest Affymetrix annotation files from Thermo Fisher database, each probe set ID was annotated to gene ID, thereafter, probe IDs that annotated to different gene IDs or did not annotate to any gene ID were excluded. When multiple probe set IDs were mapped to the same gene, probes with the highest coefficient variation (CV) over all samples were selected. The CV for each probe was computed by dividing the standard deviation of expression values among all sample population by the mean expression value.

2.3 Determining the molecular subtypes of breast cancer patients
Gene expression classifier (PAM50) of the breast cancer subtypes was applied to assign BC patients to one of the intrinsic molecular subtypes: luminal A, luminal B, HER2-enriched, triple-negative/basal-like, and normal-like.\textsuperscript{36,37} The subtype classification of each patient was determined based on the closeness between the average expression profile of 50 genes in each subtype centroid and the corresponding gene expression pattern of patient tumor, where the distances measured utilizing Spearman's rank correlation.\textsuperscript{36} From \textit{genefu} package in R, \textit{intrinsic.cluster.predict} function with pam50\textsuperscript{38} was utilized to eliminate samples with HER2-enriched, triple-negative/basal-like, and normal-like subtypes (i.e., non ER+).

### 2.4 Single-Sample Gene Set Enrichment Analysis (ssGSEA)

For single-sample analysis, gene expression values for each gene were transformed into standardized scores (i.e., z-scores) in order to bring the expression level into a common scale across all samples.\textsuperscript{39,40} Z-score for each gene was computed by subtracting the average intensity of the gene from the intensity of this gene in each sample and dividing it by the standard deviation (SD) across all samples.\textsuperscript{39} In this way, each gene’s mean is standardized to 0 and standard deviation to 1. Ranked list of z-scored for each gene for a given sample then defines a single-sample signature, utilized for further pathway enrichment analysis.

For pathway enrichment analysis, we utilized Reactome,\textsuperscript{41} BioCarta\textsuperscript{42} and KEGG\textsuperscript{43} databases, which contains 833 biological pathways, and implemented single-sample GSEA (i.e., ssGSEA\textsuperscript{44,45}), where each single-sample signature was used as a reference,
and each pathway (i.e., genes from each pathway) was used as a query set. The GSEA normalized enrichment scores (NESs), and p-values were assessed utilizing 1,000 gene permutations. NES for each of the 833 pathways (i.e., also referred to as pathway activity levels) indicated how much each pathway is overrepresented in each single-sample signature. In particular, the positive NES would indicate a pathways enrichment in the top of the rank-ordered list (i.e., overexpressed part) of the signature and the negative NES would indicate pathway enrichment in the bottom of the rank ordered list (i.e., underexpressed part) of the signature.

2.5 Associating the activity levels of molecular pathways with therapeutic response

The activity levels of each pathway (i.e., NES) were then associated with tamoxifen response using Cox proportional hazards model, adjusted for common covariates, such as age, tumor grade, tumor size, lymph node status, and PR status. For this, we utilized R coxph function from survival package. To establish a robust threshold which should be utilized to select most significantly associated pathways, we evaluated predictive ability of the pathways as a group (starting from the most significant pathway and then adding the next most significant pathway, one at a time). Thus, the groups of pathways that were evaluated were (i) Pathway 1; (ii) Pathways 1 and 2; (iii) Pathways 1, 2, and 3; etc. until all pathways were utilized. We then evaluated predicted ability of each group and recorded them (see Results). The cutoff point was determined as the one, where the addition of a pathway would not benefit an overall predictive ability of the group.
Furthermore, given that many of the 833 pathways exhibit parent-child relationships are heavily overlapping, we examined all final pathways that had the above relationships and if such dispute occurred, we prioritized pathways with higher association with tamoxifen response.

2.6 Clinical validation in independent patient cohorts

For validation studies, the activity levels of the five candidate pathways were used to stratify patients based on the risk of relapse due to treatment resistance in independent Test cohorts. Patient cohorts were subjected to t-SNE clustering, a widely-utilized dimensionality reduction technique,\(^48\) using all pairs of high-dimensional (i.e., 5-dimensions in this study) points.\(^49,50\) In fact, t-SNE reduces high-dimensional dataset (i.e., 5-dimensional) in a low-dimensional (i.e., 2-dimensional) space and successfully distinguishes groups of patients that have similar pathway activity levels. Subsequently, k-means clustering\(^51\) was utilized on t-SNE-derived the low-dimensional (i.e., 2-dimensional) space to obtain two groups of patients with distinct pathway activity patterns,\(^49,50\) using \textit{kmeans} function in R.\(^52\)

The ability of the activity levels of the 5 molecular pathways to efficiently distinguish patient clusters was determined through receiver operating characteristics (ROC) analysis\(^53\) on multiple (i.e., multivariable) logistic regression model, where normalized enrichment scores of 5 pathways were used as input parameters (i.e., independent/predictor variables) and patient clusters were utilized as a dependent/response variable. ROC curves were assessed using the area under the curve (AUC),\(^54\) where AUC
score of 0.5 indicates a random predictor. The logistic regression analysis was conducted using \textit{glm}\textsuperscript{55} function, and ROC analysis was performed using \textit{pROC}\textsuperscript{56} and \textit{ggplot2} packages in R.

Differences in therapeutic response between the patient groups were evaluated through Kaplan-Meier treatment-related survival analysis\textsuperscript{57} and Cox proportional hazards model using \textit{survival} and \textit{survminer} packages \textsuperscript{46} in R. Log-rank p-value was utilized to assess the statistical significance of the Kaplan-Meier survival analysis and Wald p-value and hazard ratio were utilized for multivariable Cox proportional hazards model through \textit{survdiff} and \textit{coxph} functions from \textit{survival} package.

To estimate the predictive accuracy of our model and obtain a more accurate indication of how well our finding behaves toward a new incoming patient, we conducted Leave-one-out cross-validation (LOOCV).\textsuperscript{58} In this method, one patient is “excluded/eliminated” and the rest of the patients are utilized for training purposes to the regression model. After that, a removed patient is assumed to be a new incoming patient and is assigned a risk of developing tamoxifen resistance. This process is repeated for each patient within a given dataset. LOOCV was implemented for multiple logistic regression model, where patient clusters membership was used as a response variable and normalized enrichment scores of our candidate pathways were utilized as input parameters. The logistic regression analysis was performed using \textit{glm}\textsuperscript{55} function, and LOOCV analysis was prepared using \textit{cv.glm} function from \textit{boot} package in R.

To evaluate non-random predictive ability of the defined 5 candidate pathways, we used random model prepared by selecting 5 biological pathways at random, thereby we investigated the statistical significance of our finding by comparing the ability of the
candidate pathways to predict tamoxifen response to random equally-sized pathways. In
details, 5 pathways were randomly chosen 1000 times from a total of 833 molecular
pathways produced from Reactome, BioCarta and KEGG databases, and subsequently
Kaplan-Meier survival analysis was utilized to evaluate the ability of random selected 5
pathways to predict therapeutic response. The empirical p-value for the significance was
estimated as the number of times predictive ability of 5 random pathways reached or
exceeded performance of our candidate 5 pathways.

2.7 Comparative analysis to other commonly utilized approaches

To assess the superiority of our approach over other commonly used techniques,
we compared its performance to (i) extreme-responder analysis;\textsuperscript{27} (ii) SVM;\textsuperscript{30} and (iii)
PRES random forest.\textsuperscript{31} In each case, we utilized Training cohort for model training and
Test cohort 1 for model validation. We compared groups of patients with poor and
favorable response to tamoxifen in the Training cohort by selecting: patients that
experienced events within 1 year of tamoxifen administration (i.e., non-responders, \( n = 4 \));
and patients that did not experience any relapses for more than 9 years (i.e., responders, \( n = 4 \)) to define a differential expression signature of tamoxifen response (i.e., through two-
sample two-tailed Welch t-test\textsuperscript{59} through \texttt{t.test} function in R). For Epsi et al method, we
then subjected the differential expression signature to pathway enrichment analysis, where
this signature was used as a reference and groups of genes from each pathway was used as
a query gene set, and treated most significant pathways as candidate pathway markers. For
SVM and PRES random forest, we subjected the differential expression signature (i.e.,
based on the proposed significance level) to the model training using Training cohort. The
SVM analysis was performed using `svm` function from `e1071` package, and PRES random forest analysis was prepared using `train` function from `caret` package in R. Predictive ability of the identified predictions was evaluated using Cox proportional hazards model through `survival` and `survminer` packages in R.

### 2.8 Pathway activity read-outs

To determine read-out genes of pathway’s activity, we examined genes inside each molecular pathway, which were (i) changed on expression levels (i.e., leading edge genes from the single sample pathway enrichment output); (ii) correlated with pathway activity outputs (i.e., correlation analysis between a leading edge gene and NESs across all patients) through Spearman correlation using `cor.test` function in R; and (iii) combined with tamoxifen response through univariable Cox proportional hazards model using adjusted hazard p-value (i.e., through `coxph` function). Finally, the adjusted hazard p-values were associated with Spearman correlation p-values to select the candidate genes.

### 2.9 Statistical analysis

Statistical analysis was performed using R studio version 3.5.1 for statistical computing. For single-sample analysis, data were z-scored on individual gene level. For this, the mean and standard deviation was first estimated for each gene across all samples in the dataset. Subsequently, z-score for each gene was defined as the difference between its own intensity value and the mean of that gene across the samples and divided by the standard deviation for that gene. The ranked list of z-scores for each gene in a sample then
defined single-sample signature. Pathway activity levels were estimated as Normalized Enrichment Scored (NESs) from the Gene Set Enrichment Analysis (GSEA), where NESs and p-values were estimated using 1,000 gene permutations. Cox proportional hazards model was utilized to associate pathway activity levels with treatment-related relapse-free survival (tRFS). When adjusting for common covariates multivariable Cox proportional hazards model was utilized, where its significance was reported using hazards ratio, hazards p-value, and Wald test. Kaplan-Meier survival analysis was utilized to estimate difference in treatment-related survival between two groups of patients, with log-rank p-value used to estimate significance.

### 2.10 Data availability

Data utilized for Training and Testing and their clinical characteristics are freely available from GEO repository GSE6532.
Chapter III

RESULTS

3.1 Overview

We present a genome-wide pathway-centric computational analysis to identify molecular pathways predictive of risk of resistance to tamoxifen in ER+ breast cancer patients. Our approach has the following steps:

Training phase (Figure 2A): (i) activity levels of biological pathways is estimated in each ER+ breast cancer patient (across a wide spectrum of responses, present in a clinical setting) that received adjuvant tamoxifen (Figure 1, Table 2); (ii) these pathway activity levels are then associated with tamoxifen treatment response across all patients, adjusted for common covariates;

Testing phase (Figure 2B): (iii) pathways that are significantly associated with the risk of tamoxifen resistance are then subjected to clinical validation analysis in independent patient cohorts (Figure 1, Table 2), for their ability to predict tamoxifen resistance for new incoming patients; (iv) finally, ability of the candidate pathways to predict the risk of tamoxifen resistance is compared to other known gene signatures of resistance and overall disease aggressiveness, alongside comparison to other methods.
Figure 2.

**A** Discovery/Training

- Patient cohort with known treatment response

- Association of treatment response with pathway activity
  - Poor response
  - Good response
  - Pathways active in poor response
  - Pathways active in good response
  - Active
  - Repressed

- Candidate pathways as markers of treatment response
  - Pathway 1
  - Pathway 2
  - Pathway 3
  - Pathway 4

**B** Testing/Validation

- Molecular profiling of patients from an independent cohort

- Patient classification based on candidate pathway activity profiles
  - Pathway 1
  - Pathway 2
  - Pathway 3
  - Pathway 4

- Clinical validation
  - ROC analysis
  - Kaplan-Meier survival
  - Cross-validation

Schematic representation of the pathway-centric approach. (A) Training phase: identification of molecular pathways of tamoxifen resistance. (B) Testing phase: clinical validation of identified candidate pathways and multi-modal prediction evaluation.
3.2 Training phase: identifying molecular pathways that govern primary tamoxifen resistance

To accurately define therapeutic response to tamoxifen in ER+ breast cancer patients, we carefully selected gene expression profiles for the Training cohort (Loi et al., KIT-GSE6532) of primary ER+ breast tumors collected through surgery, not subjected to any neoadjuvant (i.e., prior to sample collection) treatment, and administered adjuvant (i.e., post-operative) 5-year long tamoxifen administration, with available clinical follow-up data ($n = 57$) (Figure 1, Table 2).

To avoid inconsistencies in BC classification, we subjected patient profiles of the Training cohort to a 50-gene Prediction Analysis of Microarrays panel (PAM50) classification. PAM50 classification categorized BC patients from the Training cohort into the five intrinsic molecular subtypes: luminal A, luminal B, human epidermal growth factor receptor 2 (HER2)-enriched, triple-negative/basal-like, and normal-like, known to differ in their clinical outcomes and therapy choice. ER+ BC, which is the phenotype of interest in our study, is contained within the luminal A and luminal B subtypes and is excluded from HER2-enriched, triple-negative/basal-like, and normal-like subtypes (Table 2). Out of 57 post-operative tamoxifen-treated patients, 4 patients were classified as HER2-enriched, basal-like, or normal-like, and thus were excluded from further analysis.

Our objective was to evaluate tamoxifen response across all 53 patient samples (on the individual-patient level) and associate them with changes in biological pathway activities (Figure 3A). In order to be able to evaluate each patient sample individually, we scaled (i.e., z-scored, see Methods) gene expression profiles on individual gene levels so that each gene had mean 0 and standard deviation 1 over all samples in the Training
The list of genes ranked by their z-scores in each sample then defined an individual-patient signature. We then utilized each individual-patient signature to evaluate activity levels of biological pathways using single-sample Gene Set Enrichment Analysis (ssGSEA),\textsuperscript{44,45} where pathways were obtained from Reactome,\textsuperscript{41} BioCarta\textsuperscript{42} and KEGG\textsuperscript{43} databases, corresponding to 833 pathways. For this analysis, each patient signature was used as a reference and each pathway as a query gene set. Activity levels of biological pathways were defined by their enrichment in each patient signature, mathematically represented by the Normalized Enrichment Scores (NES) from the GSEA analysis, where positive NES corresponds to enrichment in the over-expressed part of the signature and negative NES corresponds to enrichment in the under-expressed part of the signature (Figure 2A, Figure 3A).
Training phase: pathway-centric approach identifies five biological pathways that govern tamoxifen response. (A) Schematic representation of the Testing phase of our approach: (left) patient molecular profiles are collected and analyzed; (middle) pathway activities are estimated in each patient using single-patient pathway enrichment analysis; (right) pathway activities are associated with response to tamoxifen using Cox proportional hazards modeling and are adjusted to common co-variates, including age, tumor grade, tumor size (> 2 cm vs ≤ 2 cm), lymph node status, and PR status. (B) Graphical illustration of tamoxifen-related treatment response or follow-up. Time to event (top): time interval between tamoxifen administration and earliest relapse is indicated by green line. Time to follow-up (bottom): time interval between tamoxifen administration and latest follow-up date is indicated by brown line (no tamoxifen-related events observed). (C) Heatmap representation of the pathway activity levels (i.e., NES) and their association with time to tamoxifen-related relapse or follow-up, in the Training cohort. Green line marks the group of patients with tamoxifen-related relapse, sorted from the shortest to the longest time to relapse. Brown line marks the group of patients with follow-up and without disease relapse until the latest follow-up, sorted from the shortest to longest time to follow-up.
Next essential step in our analysis was to associate changes in pathway activity levels to tamoxifen treatment response. In general, we defined treatment-related relapse-free survival (tRFS) as the interval between tamoxifen administration (which occurred immediately after surgery) and the earliest relapse (defined as local, regional, or distant metastasis) or the latest follow-up (these patients did not develop an event until their latest follow-up). When a patient had a relapse during or after the therapy administration, time to therapy related relapse was defined from therapy start to the earliest relapse (Figure 3B, top schematics, green line). When a patient never experienced a relapse, therapy-related relapse-free survival was measured from therapy start to the latest follow-up (Figure 3B, bottom schematics, brown line). In this dataset, 41.5% of patients experienced tamoxifen-related events (i.e., relapse), making it ideally suited for training purposes.

To estimate association between the activity levels of the biological pathways and tRFS across a wide spectrum of tamoxifen response (taking into account a heterogeneity of response to tamoxifen, present in a clinical setting), we utilized Cox proportional hazards model,\textsuperscript{46} ideally suited when time to event or follow-up is available. The Cox proportional hazards model was estimated between each pathway activity level (i.e., NESs, independent/predictor variable) and tamoxifen tRFS (i.e., dependent/response variable) across all 53 patients in the Training cohort. Furthermore, to account for the effect of other factors, this analysis was adjusted for commonly utilized covariates, as suggested in,\textsuperscript{63} such as age, tumor grade, tumor size (\(> 2 \text{ cm}\) vs \(\leq 2 \text{ cm}\)), lymph node status, and PR status (note that decreased PR levels are associated with increased HER2 signaling\textsuperscript{16}) (Figure 3A). Such analysis identified five molecular pathways (Figure 3C, Table 3), most significantly associated with response to tamoxifen (hazard p-value \(\leq 0.00075\), Figure 4A-B, see
Methods), including Retrograde Neurotrophin Signalling, Loss of NLP from Mitotic Centrosomes, RNA Polymerase III Transcription Initiation from Type 2 Promoter, EIF2 pathway, and Valine Leucine and Isoleucine Biosynthesis, adjusting for parent-child relationships inherent in pathway databases (see Methods, Figure 5).

Table 3.

Five molecular candidate pathways and their corresponding significance levels.

| Pathway Names                                           | Adjusted Hazard ratio (95% CI) | Adjusted Hazard p-value |
|---------------------------------------------------------|--------------------------------|-------------------------|
| REACTOME: RETROGRADE NEUROTROPHIN SIGNALLING            | 2.31(1.48-3.60)                | 0.00021                 |
| REACTOME: LOSS OF NLP FROM MITOTIC CENTROSOMES          | 1.73(1.28-2.33)                | 0.00029                 |
| REACTOME: RNA POLYMERASE III TRANSCRIPTION INITIATION FROM TYPE 2 PROMOTER | 1.97(1.33-2.91)                | 0.0006                  |
| BIOCARTA: EIF2 PATHWAY                                  | 1.84(1.30-2.59)                | 0.00053                 |
| KEGG: VALINE LEUCINE AND ISOLEUCINE BIOSYNTHESIS        | 1.78(1.27-2.50)                | 0.00075                 |

Note: CI: confidence intervals.
Comprehensive threshold analysis identifies pathway significance level. Threshold analysis in the Training cohort utilizing Cox proportional hazards model on the group of pathways (starting from the most significant pathways and adding the next most significant pathway, one at a time). Cutoff point was determined as a point on the graph when adding any additional pathway would not improve the model significance. Adjusted hazard p-value (A) and adjusted Cox Wald p-value (B) are used as threshold-deciding criteria.
Graphical representation of the five candidate pathways and their significantly contributing genes. Network-based representation of the five candidate pathways. Selected genes shown (brown nodes) correspond to genes that contribute to significant enrichment of each pathway in the patient single-sample signatures. Node sizes represent number of times each gene appears in the leading edge in the single-sample pathway enrichment analysis (i.e., indicating significant changes in activity of this pathway across Training cohort).
3.3 Testing phase: clinical validation in independent patient cohorts

The next essential step in our analysis was to evaluate the ability of five candidate pathways to predict treatment response to tamoxifen in independent non-overlapping clinical cohorts. For this, we utilized two patient cohorts for testing/validation purposes: (i) Test cohort 1\textsuperscript{29} (GYYT-GSE6532, \(n = 70\)) of primary breast tumors obtained at surgery, from patients that did not receive any neoadjuvant treatment and received only adjuvant tamoxifen, with 28.78\% of patients having tamoxifen-related events (Table 2); and (ii) Test cohort 2\textsuperscript{29} (OXFT-GSE6532, \(n = 77\)) of primary breast tumors obtained at surgery, from patients that did not receive any neoadjuvant treatment and received only adjuvant tamoxifen, with 25.97\% of patients with tamoxifen-related events. Both Test cohorts had clinical characteristics, neoadjuvant, and adjuvant conditions comparable to the Training cohort (Table 2). Similar to the analysis done on the Training cohort, we performed PAM50 classification on the two Test cohorts, eliminating 4 patients from Test cohort 1 and keeping all patients for Test cohort 2.

Our main objective was to investigate if activity levels of the five candidate pathways could predict risk of resistance to tamoxifen in two independent Test cohorts. For this, we estimated activity levels for five candidate pathways in each patient in the Test cohorts (similarly to Training cohorts, see Methods) and subjected patients to t-distributed Stochastic Neighbor Embedding (t-SNE) clustering\textsuperscript{48} as suggested in\textsuperscript{49} for investigation of samples relationships. T-SNE analysis, which displays five-dimensional dataset in a two-dimensional space, stratified patients into two groups based on their pathway activity levels. The low-dimensional output (i.e., 2-dimensional) of t-SNE were then subjected to the k-means clustering\textsuperscript{51} to correctly assign group membership (Figure 6A for Test cohort
1 and Figure 6C for Test cohort 2) one group with increased pathways’ activities (orange) and one group with decreased pathways’ activities (turquoise), mimicking the relationship that was observed in the Training cohort (Figure 3C). We confirmed the strength of group separation through Receiver Operating Characteristic (ROC) analysis\textsuperscript{53} using multiple logistic regression model (Figure 7A-B), where normalized enrichment scores of 5 pathways were used as input parameters (i.e., independent/predictor variables) and selected patient groups were utilized as a dependent/response variable. The efficiency of ROC analysis was estimated using area under the curve (AUC),\textsuperscript{54} where AUC of 0.5 denotes a random predictor and AUC score of 1 denotes a perfect predictor (i.e., full separation of the patient groups). This analysis confirmed that the activity levels of the five candidate pathways can be effectively used for classifying patients into distinct groups (Test cohort 1, AUC = 0.929; Test cohort 2, AUC = 0.867).
Figure 6.
The five candidate pathways predict patients at risk of tamoxifen resistance in independent patient cohorts. (A, C) T-SNE and subsequent k-means clustering of Test cohort 1 (A) and Test cohort 2 (C) based on activity levels of the five candidate pathways demonstrates patient separation into two groups: orange group (with overall increased activity levels of the five candidate pathways) and turquoise group (with overall decreased activity levels of the five candidate pathways). (B, D) Kaplan-Meier treatment-related survival analysis comparing two patient groups in Test cohort 1 (B) and in Test cohort 2 (D). Log-rank p-values and adjusted hazard ratios are indicated. (E, F) Leave-one-out cross-validation (LOOCV) correctly identified patients with poor response to tamoxifen (orange) and patients with favorable response to tamoxifen (turquoise) in Test cohort 1 (E) and Test cohort 2 (F). Accuracy values (%) are indicated. (G) Random model to assess the ability of the 5 candidate pathways selected at random to differentiate samples into groups with various tamoxifen response. The significance of the predictive ability of our defined 5 molecular pathways is shown by the distributions of log-rank p-values from the random model.

**Figure 7.**

ROC analysis demonstrated significant separation of patient groups based on activity levels of the five candidate pathways. ROC analysis to show significance of the separation between patient groups in Figure 6 A, C. Area under the curve (AUC) is reported.

To assess if these patient groups significantly differ in their tamoxifen response, we analyzed therapy-related relapse-free survivals between the groups using Kaplan-Meier survival analysis \(^{57}\) and Cox proportional hazards model \(^{46}\), which demonstrated that the
identified patient groups had a significant difference in their response to tamoxifen (Test cohort 1, log-rank p-value = 0.02, Figure 6B; Test cohort 2, log-rank p-value = 0.01, Figure 6D). We have also adjusted these analyses for common covariates65 (i.e., age, tumor grade, tumor size, lymph node status, and PR status), demonstrating that these covariates did not significantly impact the predictive ability of our findings (Test cohort 1, adjusted hazard ratio = 3.11, adjusted hazard p-value = 0.044, 95% confidence interval CI: = 1.03-9.396, Figure 6B; Test cohort 2, adjusted hazard ratio = 4.24, adjusted hazard p-value = 0.012, CI: 1.3708 - 13.120, Figure 6D).

Further, we evaluated predictive accuracy of our model in the two test cohorts using Leave-one-out cross-validation (LOOCV), which simulates a situation when a new incoming patient needs to be evaluated for her risks of developing resistance to tamoxifen. In particular, in LOOCV, one patient is “removed”, and the model is trained on the remaining patients, followed by the prediction of risk of resistance for the removed patient. The process is repeated for each patient. Using this analysis, we demonstrated the accurate performance of our model in predicting poor and favorable tamoxifen response for new incoming patients (Test cohort 1, accuracy for LOOCV = 85.8%, Figure 6E; Test cohort 2, accuracy for LOOCV = 82.5%, Figure 6F). Finally, to evaluate if any set of five pathways selected at random could classify patients based on tamoxifen response, we have performed random model analysis, where selected five pathways at random 1,000 times and evaluated their predictive ability using Kaplan-Meier survival analysis, as above. The empirical p-value for the random model was estimated as the number of times log-rank p-values of five pathways chosen at random reached or exceeded the output of our 5 candidate pathways, which confirmed significant non-randomness of our candidate pathways predictive ability
(Test cohort 1, random model p-value = 0.031, Figure 6G). Taken together, these findings indicate that the five-candidate pathway signature could successfully predict patients at risk of tamoxifen resistance in independent patient cohorts.

3.4 Comprehensive comparison of tamoxifen response and overall disease aggressiveness

A fundamental question in studying therapeutic response lies in its comparison to and differentiation from overall disease aggressiveness. Our comprehensive investigation of this question was four-fold: (i) we identified pathways implicated in disease aggressiveness and compared their overlap with the candidate five pathways of tamoxifen response; (ii) we evaluated if the five candidate pathways can predict breast cancer aggressiveness in an independent (negative control) cohort; (iii) we evaluated the ability of the five candidate pathways to predict tamoxifen response based on PR status (known indicator of breast cancer aggressiveness), age categories (as patients aged 50 years or older shows poorer relative survival rates than younger patients 64) and luminal subtypes (as luminal B type have a poorer prognosis than luminal A type 65); and (iv) we evaluated if known published signatures of disease aggressiveness could predict response to tamoxifen.

First, to examine if our 5 candidate pathways overlap with pathways implicated in disease aggressiveness, we developed treatment-free prognostic pathway signature using a patient cohort that received surgery only (KIU-GSE6532, n = 51, negative control cohort).29 Out of 51 surgery-treated patients, 4 patients were removed, based on the PAM50 classification. We further applied our single-sample pathway-based discovery approach (as in the Training phase) and associated them to the RFS, which identified 3 pathways of
aggressiveness (see Methods) that showed no overlap with the five candidate pathways, signifying that none of our candidates are involved in cancer severity and are indeed specific to tamoxifen response.

Figure 8.

Five candidate pathways do not predict overall disease aggressiveness. (A) T-SNE and subsequent k-means clustering based on the activity levels of the five candidate pathways in the negative control cohort. (B) Kaplan-Meier survival analysis on negative control cohort confirms that the five candidate pathways do not predict disease aggressiveness. Log-rank p-value and hazard ratio are indicated.
Stratified analysis demonstrates that predictive ability of the five candidate pathways is not dependent on the PR status. Patients in Test cohort 1 were stratified based on their progesterone PR status: Progesterone-positive (A, B) and Progesterone-negative (C, D). T-SNE with subsequent k-means clustering on PR+ (A) and PR- (C) patient subgroups. Kaplan-Meier survival analysis for PR+ (B) and PR- (D) patient groups. Log-rank p-values are indicated.
Secondly, we evaluated if the five candidate pathways could separate patients based on overall disease aggressiveness. For this, we evaluated predictive ability of the five candidate pathways on the BC patient cohort that did not receive any treatment after surgery (negative control cohort, as above). We subjected the dataset to the single-sample pathway enrichment analysis (for the five candidate pathways, similarly to Test cohorts analysis). T-SNE clustering (Figure 8A) and subsequent Kaplan-Meier survival analysis (Figure 8B) on this cohort demonstrated that the five pathways do not separate patients based on their disease aggressiveness (hazard ratio = 1.2, log-rank p-value = 0.7, RFS was considered as a clinical endpoint), but rather specific for tamoxifen response. We have also examined the effect of covariates (i.e., age, tumor grade, tumor size, and PR status), on disease progression in this setting and demonstrated that our candidate pathways remain insignificant, with tumor size significantly contributing to the disease progression (adjusted hazard p-value = 0.0307).

Third, given that the PR receptor status (which also reflects HER2 signaling) is a known indicator of breast cancer aggressiveness, we performed a stratified Kaplan-Meier analysis on Test cohort 1 (for which this information was available). For this, we divided Test cohort 1 into two groups: one with PR-positive status and one with PR-negative status. We then subjected both cohorts separately to t-SNE clustering, which have demonstrated that the five candidate pathways separated each cohort into patient sub-groups with high and low levels of pathway activities (Figure 9A for patients with PR-positive tumors and Figure 9C for patients with PR-negative tumors). Subsequent Kaplan-Meier survival analysis (Figure 9B and Figure 9D, respectively) showed that these patient-subgroups significantly differ in their response to treatment (patients with PR-positive tumors, log-
rank p-value = 0.02, Figure 9B; patients with PR-negative tumors, log-rank p-value = 0.01, Figure 9D), demonstrating that our five candidate pathways are able to predict patients at risk of tamoxifen resistance regardless of the PR-status.

**Figure 10.**

Stratified analysis demonstrates that predictive ability of the five candidate pathways does not depend on the age groups and luminal subtypes. Stratified Kaplan-Meier survival analysis was conducted using different age groups in Test cohort 2 (A, B) and luminal subtypes in Test cohort 1 (C, D).
In fact, different age categories and luminal subtypes are also known indicators of worse prognosis in breast cancer. Thus, we conducted stratified Kaplan-Meier analysis (as above); where we stratified Test cohort 2 into patient groups based on age (< 50 years and >=50 years), and Test cohort 1 into patient groups based on luminal subtypes (luminal A and luminal B). Kaplan-Meier survival analysis showed clear separation between patient groups (Figure 10 A-D). Our analysis demonstrated that the predictive ability of these candidate pathways does not depend on known characteristics of breast cancer aggressiveness.

Figure 11.

Five candidate pathways are not affected by overall disease aggressiveness. Multivariable Cox proportional hazards model representing analysis for five candidate pathways adjusted for various prognostic signatures in breast cancer, including Wang et al. (76 prognostic markers, with 57 present on U133 Plus 2.0) and van’t Veer et al. (70 prognostic markers, with 53 present on U133 Plus 2.0). Adjusted hazard p-values are reported.
Finally, to demonstrate that the predictive ability of the five candidate pathways is not affected by other known markers of disease aggressiveness, we investigated if commonly known gene-based prognostic signatures can predict tamoxifen response or affect predictive ability of the candidate five pathways. For this, we gathered several known signatures of overall BC aggressiveness (i.e., prognostic signatures), including Wang et al. signature\(^66\) (76 prognostic markers, with 57 present on U133 Plus 2.0) and van’t Veer et al. signature\(^67\) (70 prognostic markers, with 53 present on U133 Plus 2.0) and subjected them to adjusted multivariable Cox proportional hazards model, alongside the five candidate pathway signature, in the Test cohort 1. This analysis confirmed that the prognostic signatures were not predictive of tamoxifen response and did not impact predictive ability of the five candidate pathways (adjusted hazard p-value = 0.03, Figure 11). Taken together, these findings indicate that our five-pathway signature of tamoxifen response is not indicative of overall breast cancer aggressiveness and is indeed specific to response to tamoxifen.

3.5 Comparative analysis to commonly utilized methods and known signatures of tamoxifen response

To evaluate predictive advantages of the five candidate pathways, we took a comprehensive approach and first (i) compared the predictive ability of the five candidate pathways to predictions from other commonly used methods, including approaches based on extreme-responder analysis (i.e., tails of the distribution), support vector machine (SVM), and random forest; and second (ii) assessed if the predictive ability of the five candidate pathways outperforms other known signatures of tamoxifen response.
Predictive ability of the five candidate pathways outperforms markers from other methods and known signatures of tamoxifen response. (A, B) Comparison of the predictive ability of the five candidate pathways (blue) to the candidate identified by other approaches, including Epsi et al. extreme-responder analysis (green), Zhong et al. SVM-based method (brown) and Yu et al. PRES random forest-based method (pink), through unadjusted (A) and adjusted for common covariates (B) Cox proportional hazards model. P-values for unadjusted and adjusted hazard ratios are indicated. (C) Multivariable Cox proportional hazards model representing analysis for the five candidate pathways adjusted for different predictive signatures of tamoxifen response, including Men et al. (10 predictive markers, with 9 present on U133 Plus 2.0), Paik et al. (Oncotype DX, 21 predictive markers), and Ma et al. (2 predictive markers). Adjusted hazard p-values are indicated.

First, we compared predictive ability of the five candidate pathways to predictions from other commonly utilized methods, such as (i) Epsi et al.\textsuperscript{27} method, which utilized extreme-responder analysis, using tails of the treatment response distribution to define a treatment response signature; (ii) Zhong et al.\textsuperscript{30} method, which used Support Vector Machine approach as a base; and (iii) Yu et al.\textsuperscript{31} method, also referred to as Personalized REgimen Selection (PRES), which used random forest approach as a base (see Methods). To assure that all methods are comparable to our pathway-centric method, we trained Epsi et al., Zhong et al., and Yu et al. methods on the Training cohort, with each producing a list of predictions (112 predictions for Epsi et al.; 5 predictions for Zhong et al.; and 3
predictions for Yu et al.). We then followed by validating these predictions on the Test cohort 1, similarly to our pathway-centric method. Such analysis demonstrated that the five candidate pathways outperform all three methods in their ability to predict the risk of tamoxifen treatment resistance (Figure 12A: five candidate pathways, hazard ratio = 2.91, hazard p-value = 0.031; Epsi et al., hazard ratio = 2.79, hazard p-value = 0.038; Zhong et al., hazard ratio = 2.53, hazard p-value = 0.063; Yu et al., hazard ratio = 2.48, hazard p-value = 0.058). Furthermore, we adjusted these analyses for the effect of common covariates (similarly to our original training phase), including age, tumor grade, tumor size, lymph node status and PR status and re-confirmed that the five candidate pathways retain their significant predictive ability and outperform the other methods (Figure 12B: five candidate pathways, adjusted hazard ratio = 3.11, adjusted hazard p-value = 0.044; Epsi et al., adjusted hazard ratio = 2.48, adjusted hazard p-value = 0.076; Zhong et al., adjusted hazard ratio = 2.96, adjusted hazard p-value = 0.05; Yu et al., adjusted hazard ratio = 2.81, adjusted hazard p-value = 0.054).

Finally, to confirm that the predictive ability of the five candidate pathways outperforms other known signatures in their ability to predict tamoxifen treatment response, we selected known signature of tamoxifen response (i.e., predictive signatures), such as (i) Men et al.\textsuperscript{18} (10 predictive markers, with 9 present on U133 Plus 2.0); (ii) Paik et al.\textsuperscript{19} (also now as Oncotype DX, 21 predictive markers); and (iii) Ma et al.\textsuperscript{20} (2 predictive markers) (Figure 12C) and used them in adjusted multivariable Cox proportional hazards model, alongside the five candidate pathway signature, utilizing Test cohort 1, as above. This analysis demonstrated that the additional predictive signatures do not significantly affect the ability of the five candidate pathways to predict the risk of tamoxifen resistance.
(Figure 12C, adjusted hazards p-value = 0.03). Taken together, these results demonstrate that the five-candidate pathway signature can be utilized to predict patients at risk of developing resistance to tamoxifen in a clinical setting and build a foundation for personalized therapeutic advice for patients with ER+ breast cancer.

Figure 13.

The five read-out genes predict patients at risk of tamoxifen resistance in independent patient cohort. (A) T-SNE and subsequent k-means clustering of Test cohort 1 based on gene expression of the five read-out genes demonstrates patient separation into two groups. (B) Kaplan-Meier treatment-related survival analysis comparing two patient groups in Test cohort 1. Log-rank p-value and hazard ratio are indicated.
Table 4.

Five read-out genes and their corresponding significance levels.

| Read-out genes | Spearman correlation p-value | Adjusted Hazard p-value |
|----------------|-----------------------------|-------------------------|
| AP2S1          | 0.0000278                   | 0.0185                  |
| CDC2           | 0.00000035                  | 0.0063                  |
| GTF3C3         | 0.00000004                  | 0.00133                 |
| EIF2AK3        | 0.00000542                  | 0.019                   |
| LARS           | 0.00000013                  | 0.019                   |

3.6 Pathway activity read-outs

For this, we examined genes which could function as read-outs of activity levels of pathways that involved in treatment response. In details, we examined genes inside each molecular pathway, which were (i) changed on expression levels (i.e., from leading edge in pathway enrichment outputs); (ii) correlated with pathway activity outputs (i.e., NESs); and (iii) combined with tamoxifen response (see Methods, Table 4). This analysis defined 5 read-out genes (i.e., AP2S1, CDC2, GTF3C3, EIF2AK3, and LARS) that were significantly related to therapeutic response (Test cohort1: log-rank p-value = 0.03, hazard ratio = 3.76, Figure 13 A-B). In fact, these findings showed a high predictive ability in identifying patients at risk of resistance. We suggest that these 5 read-out genes can be utilized as biomarker of tamoxifen resistance and will be more useful in the clinic.
Chapter IV

DISCUSSION

In this study, we have demonstrated that a pathway-centric genome-side computational approach is able to uncover biological pathways, highly associated with risk of tamoxifen resistance in ER+ breast cancer patients. The important advantage of our approach is that it identifies a tightly connected group of genes - biological pathways - as opposed to individual (possibly distantly connected genes), thus (i) decreasing the chances of experimental noise present in biological experiments; (ii) improving our understanding of the mechanisms implicated in therapeutic resistance; and (iii) increasing the likelihood of identifying a functionally relevant signature, which could be utilized to study mechanisms of primary resistance and their potential therapeutic targeting. Furthermore, these biological pathways are highly associated with a wide spectrum of treatment responses (as opposed to selecting a limited category of patients for analysis), reflecting heterogeneity of response to tamoxifen present in a clinical setting. Even though this work is focused on identifying cases of resistance to tamoxifen, our method can be broadly applicable to other therapeutic interventions and cancer types.

Our computational analysis has identified five molecular pathways implicated in tamoxifen resistance, including (i) Retrograde Neurotrophin Signalling, (ii) Loss of NLP from Mitotic Centrosomes, (iii) RNA Polymerase III Transcription Initiation from Type 2 Promoter, (iv) EIF2 pathway, and (v) Valine Leucine and Isoleucine Biosynthesis. Interestingly, many of these pathways have been shown to be closely related to carcinogenic mechanisms and therapeutic response in various cancers. In particular, the
Retrograde Neurotrophin Signalling pathway is implicated in metabolic detoxification, mitosis, clathrin-mediated vesicles development, and enriched with bladder cancer predisposition loci. One of the genes from this pathway, Neurotrophic tyrosine kinase receptor type 1 (NTRK1), is a recognized oncogene frequently altered in various tumor types and its gene fusions have previously been identified in glioblastoma, colon cancer, papillary thyroid carcinoma, and non-small cell lung cancers. Clinical studies of tumor response to NTRK1 fusion-targeted therapy have indicated that this oncogene represents a treatment target in human cancer.

Ninein-like protein (NLP) (i.e., also known as NINL) is a part of the Loss of NLP from the Mitotic Centrosomes pathway. The role of human centrosomal NLP expression in breast, lung, ovarian, head and neck cancers has been widely demonstrated. The NLP gene amplification accounts for NLP overexpression in human breast and lung cancer cells. The deregulated expression of NLP in cell models leads to mitotic spindle aberrations, spindle checkpoint defects, chromosomal missegregation, cytokinesis failure, stimulation of chromosomal instability, anchorage-independent growth, and cell malignant transformation. Recently, it has been discovered that NLP co-localizes and interacts with BRCA1 at inter-phasic centrosome and thus the disruptions of BRCA1 function could affect NLP co-localization to centrosomes and induce the genomic instability. Interestingly, it has been reported that the NLP overexpression may also cause breast cancer resistance to paclitaxel chemotherapy. Furthermore, a positive correlation between expression of NLP and PLK1 (i.e., another gene implicated in the Loss of NLP from the Mitotic Centrosomes pathway) has recently been discovered, implicated in
chemoresistance, particularly to taxane agents\textsuperscript{76} and tumor growth in general, in breast cancer and other cancer types.\textsuperscript{76,77}

For the RNA Polymerase III Transcription Initiation from RNA polymerase II Promoter Sites, the global gene expression is increased in eukaryotic cells as RNA polymerase II transcribes protein-coding genes to yield mRNA, miRNA, snRNA, and snoRNA genes while RNA polymerase III transcribes the genes for 5S rRNA and tRNAs.\textsuperscript{78,79} However, in yeast (S. cerevisiae), RNA polymerase III complex has been shown to act as heterochromatin barriers,\textsuperscript{80} and any change in heterochromatin would be potentially very important for cancer development. A broad spectrum of cancer cell types has been observed to display a highly regulated and elevated level of RNA polymerase III transcript expression.\textsuperscript{81,82}

In the Eukaryotic Initiation Factor 2 (eIF2) pathway, phosphorylation of eIF2α has been shown to play a significant role in maintaining normal cellular homeostasis and regulating cell growth.\textsuperscript{83} with dysregulation of eIF2 signaling pathway stimulating the cancerous tumors transformation.\textsuperscript{84} The overexpression of eIF2α has been observed in several cancers, such as gastrointestinal cancer\textsuperscript{85} and non-Hodgkin's lymphomas\textsuperscript{86} and has been proposed as a potential therapeutic target.\textsuperscript{87}

Finally, in the Valine Leucine and Isoleucine Biosynthesis pathway, valine, leucine, and isoleucine are important branched-chain amino acids (BCAAs) for normal growth and development.\textsuperscript{88} In the BCAA catabolism pathway, the first step is transamination, catalyzed by the branched chain amino acid transferase isozymes BCATs: a mitochondrial (BCATm) and a cytosolic (BCATc) isozyme.\textsuperscript{89-91} Mitochondrial BCATm (BCAT2) expression can drive the development of pancreatic ductal adenocarcinoma under the
regulation of the mitochondrial malic enzyme. Malic enzyme 2 and malic enzyme 3 are oxidative decarboxylation that stimulate malate to pyruvate and are considered important elements during mitochondrial reactive oxygen species homeostasis and NADPH production. Studies have reported that incorporating the molecular and metabolomic examination of malic enzyme-deficient cells showed a decrease in NADPH generation and an increase in reactive oxygen species level. These alterations catalyze AMPK which consequently functions to inhibit SREBP1 and thereby suppresses its target genes including the BCAT2. BCAT2 stimulates the transfer of the amino group of BCAAs to α-ketoglutarate and thus generates glutamate, which in turns enhance de novo nucleotide synthesis. Therefore, the deficiency of the mitochondrial malic enzyme, which causes a reduction in NADPH synthesis, plays a critical role in the therapeutic strategy for the treatment of patients with complex disease.

Cytosolic BCATc (BCAT1) is overexpressed in glioblastoma, nasopharyngeal carcinoma, and cancers with elevated c-MYC. It has been recommended to consider BCAT1 as a promising target for glioblastoma and nasopharyngeal carcinoma treatments. We propose that the identified candidate pathways should be investigated for their potential use as isolated treatment targets or in combination with ER-targeting agents for ER+ breast cancer patients at risk of developing resistance to tamoxifen.

One of the limitations of our study is in the limited availability of the epigenomic profiles for our patient cohorts. In fact, DNA and histone methylation has been suggested to be responsible for inactivation of ER. Thus, further examination of the role of epigenomic modulations and their interplay with transcriptomic changes is an invaluable
next step for in-depth understanding of molecular mechanisms implicated in hormone therapy resistance.

Furthermore, miRNAs (micro-RNAs) have received substantial attention for their role in regulating pathway functionality.\(^97\) For example, miR-15a/miR-16’s deletion or down-regulation contributes to dysregulated of cell cycle in chronic lymphocytic leukemia\(^98\) and non-small cell lung cancer.\(^99\) Even though miRNA data are not available in our cohorts, we foresee the importance of miRNA analysis for further understanding mechanisms of pathway dysregulation, especially when applies to therapeutic resistance.\(^100\)-\(^102\) The presence of miRNAs in tumor-derived exosomes has recently been postulated to play important roles in facilitating metastasis, and this work suggests that exosomes containing tumor-derived miRNAs which regulate one of these five pathways may also play a role in the spread of tamoxifen resistance.\(^103\)

In addition, availability of single-cell profiles for investigation of therapeutic response has proven to be invaluable\(^104\) in understanding of therapeutic targets for complex diseases, including cancer. Thus, as such profiles become available, we foresee their immediate utilization for elucidation of mechanisms of primary and secondary therapy resistance, and we investigate the miRNAs which are known to regulate any of the 5 molecular pathways.
Chapter V

CONCLUSIONS

In conclusion, we have demonstrated that a systematic computational pathway-centric method could identify molecular pathways to predict tamoxifen resistance, including (i) Retrograde Neurotrophin Signalling, (ii) Loss of NLP from Mitotic Centrosomes, (iii) RNA Polymerase III Transcription Initiation from Type 2 Promoter, (iv) EIF2 pathway, and (v) Valine Leucine and Isoleucine Biosynthesis. We propose that our finding can be ultimately utilized to prioritize and determine (i) cases at higher risk of developing resistance to tamoxifen that should be considered for alternative treatment manipulations (for instance, alternative endocrine therapy, radiation therapy, or chemotherapy etc.) and (ii) cases who would benefit maximally from tamoxifen therapy.
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