Molecular landscape: a data integration tool for personalized medicine in Breast Cancer

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# Abstract

Estrogen receptor (ER) signaling is an important factor in breast carcinogenesis. A breast cancer patient that has more than 1% (or 10%) of ER positive cells by immunohistochemistry (IHC) usually receives standard endocrine therapy. We hypothesized that there are differences in endocrine therapy response among estrogen receptor positive patients based on the molecular estrogen signaling. We show that a continuous molecular measure of estrogen signaling is prognostic among all ER+ breast cancer patients and also among those with high ER IHC percentage (). We propose a new framework for personalized medicine to better understand the molecular biology of tumors. We introduce a novel single sample embedding technique based on RNA-seq and microarray data that integrates distinct cohorts, such as TCGA, METABRIC and SCANB. We show that the embedding, also called molecular landscape, captures the true biology of breast cancer, dividing patients based on ER status and the different molecular subtypes. The embedding works in a way that the algorithm is able to project new samples together within the three cohorts, allowing to compare them to what is publicly available and avoiding the need of specific reference sets. We demonstrate how to use the molecular landscape to ask specific questions about patients in terms of estrogen signaling in the context of the POETIC trial and what could be possible reasons that explain the response to treatment. The embedding is not dependent of tumor purity, age and other clinical variables but other pathways, such as G2M Checkpoint, Epithelial to Mesenchymal Transition (EMT) and PI3K AKT MTOR signaling drive the embedding. Lastly, we analyse specific parts of the molecular landscape to look for potential subgroups and alternative therapies, more specifically, we show that PI3K AKT MTOR signaling might be a potential target pathway to be considered in future treatments among specific subgroups on the embedding.

# Introduction

Breast cancer is the most commonly diagnosed tumor worldwide (1). It is a very heterogeneous disease that can be subdivided in different histological and molecular subtypes. Patients that are diagnosed with breast cancer are classified based on the expression of the estrogen, progesterone receptors and HER2 membrane protein. The estrogen receptor positive (ER+) breast cancer subtype is the most frequent found in clinical practice (more than 70% of the cases) (2). Tumor cells of this subtype contain the estrogen receptor whose main mechanism is the target of selective estrogen receptor modulator (SERM), selective estrogen receptor degrader (SERD) and aromatase inhibitors (AI). The degree of the expression of the protein can range from 1% to 100% positive cells according to ASCO guidelines (3). It has been shown that the patients in the low spectrum of ER positivity, from 1% to 10%, do not benefit as much as patients with 10% to 100% of ER positive cells in the tumor (4).

Transcriptomics and genomics have revolutionized cancer research. The new related tools opened the option to better understand the molecular underpinnings of cancer biology. In breast cancer, microarray technologies and RNA-sequencing have been used to aid clinicians when taking the decision of giving chemotherapy for patients, avoiding potential overtreatment of patients. Researchers have developed gene signatures that are able to assign a risk score Such risk score is based on survival data found in the literature and the higher the risk score of a patient, the higher the benefit of additional chemotherapy besides the first-line therapy, such as aromatase inhibitors. Several of these signatures are already clinically used (5–9). Moreover, some of the risk scores are already recommended in international guidelines (10). These signatures though do not provide a possible explanation on why a patient should receive additional therapy. These signatures are composed of several submodules, they are in most of the cases related to estrogen, proliferation and HER2 status. For example, the recurrence score (RS) is strongly associated with the estrogen module and in the Risk of Recurrence (ROR) signature there is a strong correlation with its proliferation module (11). An alternative to the signatures only approach is to use RNA-sequencing in the clinics (12). The SCANB team showed that it is possible to use mutational and gene expression based biomarkers within one week of tumor surgery (13–15) in the clinics. They also showed that it is possible to use RNA-seq to calculate the Prosigna risk scores and risk stratification (16). This all highlights the clinical relevance of RNA-sequencing for clinical use, holding a promising future.

There is still a lack of understanding of the patient’s tumor molecular biology. There are challenges to overcome in the pathway analysis for patients individually and how to compare patients molecularly when considering complete transcriptomics data. There is no tool that allows to integrate patients in a single sample way. Usually integration is a one step procedure and cannot be updated. The common tools used in the RNA-seq community for batch effect removal are (17–19) and new samples cannot be integrated in a straight forward manner. The only way is to re-run the procedure together with the new sample. Such procedure is not usually feasible as there are not enough samples to estimate batch effects across groups and therefore correct them.

We propose here a framework for personalized medicine and the understanding of the signaling pathways at the patient level. By developing a normalization and embedding technique, we show that it is possible to integrate publicly available molecular datasets, such as TCGA, SCAN-B, METABRIC, microarray data of some patients from the POETIC trial and also patient derived xenografts (12,20–23). By using a special normalization and principal component analysis we can combine the datasets together explore the full molecular landscape of RNA-sequencing and microarray samples together, also showing how the molecular subtypes cluster in a continuous way. The new molecular landscape allows us to add new single samples independently, showing what is its molecular subtype without the need of any other classification tool that has been validated in another dataset. It is possible to compare patients in a neighborhood of the embedding, providing a context to compare patients and better understand the biology of the individual tumor. Moreover, we propose to use the different gene sets from the hallmarks collection of the Molecular Signature Database (MSigDB) (24,25), enabling patient individualized understanding of the tumor biology.

# Methods

## Cohorts

The breast cancer cohorts TCGA, SCANB, METABRIC and POETIC (12,20–22) were used to calculate the principal component analysis (PCA) embeddings and perform the validation. The samples from TCGA, SCANB and METABRIC are representative of the different breast cancer molecular subtypes in each respective country of the study. For these three cohorts, only samples primary breast cancer were used. The POETIC trial data comprises only post-menopausal women that were diagnosed with primary breast cancer and are considered to have an estrogen receptor positive status (IHC) in the initial diagnosis.

TCGA was downloaded from Firebrowse. The version 3 of SCANB data (StringTie FPKM Gene Data unadjusted) was downloaded from Mendeley <https://data.mendeley.com/datasets/yzxtxn4nmd> (26). METABRIC was downloaded directly from cBioPortal. POETIC data was downloaded from GEO (accession code GSE105777) using the R package GEOquery. Details about the download and preprocessing steps are described in https://chronchi.github.io/transcriptomics.

## Selection of highly variable genes

In order to perform the PCA, we sub selected 1000 common and highly variable genes in the TCGA and METABRIC cohorts. For each gene and in each cohort separately, the coefficient of variation (CV) was calculated:

The 1000 genes with highest average CV were selected.

## qPCR-like normalization

Given a list of 44 stable genes across different cancers (27) and 1000 genes selected, all 1044 genes were ranked from lowest to highest expression for each sample separately and the rankings were divided by the average ranking of the 44 stable genes.

## PCA embedding

Using the normalized data as described in the qPCR-like normalization, a total of 1000 random samples (fixed seed in R) coming from TCGA and METABRIC were selected to perform the initial PCA, using PCAtools (28) in R. PCA was performed without centering and scaling since the data is already centered and scaled for all genes and samples. The embedding for new individual samples is obtained by multiplying the loadings matrix with the sample normalized expression. For samples where not all the 1044 genes are available, the normalization is performed and the missing genes are padded with 0.

## Scoring strategies

For the 4 big cohorts, TCGA, SCANB, METABRIC and POETIC, GSVA (29) was applied along with the signature (30) and the hallmark collection from the molecular signature database (24,31). Default parameters were used in the gsva function from the GSVA package.

## Average neighborhood scores

In order to calculate the posterior distribution of the average scores in each neighborhood, a linear regression with only intercept (score ~ 1) was fit using rstanarm (32) and the function stan\_glm for each pathway individually. When applying the function stan\_glm, we used four chains and a prior normal distribution with location and scale equals to . The package tidybayes (33) was used to extract the draws and put them in a tidy format.

## Survival analysis

Survival analysis was done using cox regression with the survival package from R. The variables used for adjustment were age, tumor stage and number of positive lymph nodes for TCGA and SCANB. For METABRIC the variables used were Age and the Nottingham Prognostic Index (NPI). Overall survival was performed for all three cohorts and recurrence free survival for METABRIC and SCANB. The causal model used here is that the confounders affect both the outcome and the pathway of interest. It could be that there are unmeasured confounders that are affecting the pathways and outcomes at the same time, leading to a biased result. For METABRIC the variables age and nottingham prognostic index were used for adjustment. For SCANB and TCGA the variables age, node stage and tumor stage were used for adjustment.

## Code availability

The code used to generate all the analysis is available on <https://github.com/chronchi/molecular_landscape>. To fully reproduce the images in this paper check the instructions on how to run the docker image and Rstudio session at the github repository. An online version with a website containing all the analysis can be found on <https://chronchi.github.io/molecular_landscape>.

# Results

## Estrogen receptor signaling is a clinical continuous variable

We hypothesized that estrogen receptor (ER) signaling measured by scores from bulk RNA-seq samples is prognostic among patients that received only endocrine therapy. One way to define estrogen signaling is to calculate scores derived from gene expression levels of bulk RNA-sequencing. This provides a continuous measure for estrogen signaling that can be used to predict the impact of endocrine therapy. We used three independent breast cancer molecular bulk RNA-seq datasets, TCGA, SCANB and METABRIC, (12,20,21) to calculate estrogen signaling scores. They are cohorts with over 1000 patients each that cover the whole molecular spectrum of primary breast cancer samples. The estrogen signatures HALLMARK\_ESTROGEN\_RESPONSE\_EARLY and HALLMARK\_ESTROGEN\_RESPONSE\_LATE were extracted from the molecular signature database (24) (MSigDB) and from (30). The signatures from MSigDB contain 200 genes respectively and the later signature has 18 genes that are associated with estrogen and progesterone receptor signaling. The individual scores for each patient sample are shown in [Figure 1](#fig-01) (a) for each cohort stratified by estrogen receptor status. The scores capture the differences between estrogen receptor positive and negative breast cancer as expected. There is a wide range of values in the estrogen receptor positive (ER+) subgroup among the three cohorts ([Figure 1](#fig-01) (a)).

The SCANB cohort has also estrogen receptor percentage determined by immuno histochemistry (IHC), the most commonly used measure in the clinics to decide if a patient should receive endocrine therapy or not. We show that there is a correlation between the molecular ER signaling score and ER percentage ([Figure 1](#fig-01) (b)) and that among patients with high ER IHC percentage there is a continuum of ER signaling scores.

Cox regression was used to determine the hazard ratio (HR) of the ER signaling in overall survival (OS) for TCGA, SCANB and METABRIC and recurrence free survival (RFS) for METABRIC. Each survival analysis was done independently and adjusted for available clinical variables. Tumor size and number of lymph nodes were used for TCGA and SCANB cohorts. The Nottingham prognostic index (NPI) was used for METABRIC. Age was used in all cohorts as a clinical variable for adjustment. To better understand the effect of estrogen receptor signaling in the clinics, only ER+ BC patients were used and when possible only those that received endocrine therapy (for SCANB and METABRIC). Such sampling may avoid a bias on the coefficients due to chemotherapy, as patients who receive chemotherapy tend to have a worse outcome compared to those that do not. [Figure 1](#fig-01) (c) shows the forest plots for each cohort individually when calculating the hazard ratio for the estrogen signaling signature. In all the three cases, the hazard ratio for estrogen early was below 1, with values ranging from 0.23 to 0.61. In all three cases the upper part of the confidence intervals (CI) are below 1, with TCGA having the widest CI, possibly due to the fact we could not subset based on endocrine therapy, only by molecular subtype. This shows the continuous aspect of estrogen receptor signaling. For both estrogen pathways of the hallmark collection of gene sets, hazard ratios were below 1 in all cases (Table S1), with a HR below 1 meaning that the higher the score the better the chances of survival for a patient.

Given that the ER percentage could have an impact in the previous results, we only selected patients that have high ER IHC percentage (above 90%) and performed survival analysis once again on the SCANB cohort. The HR for ER percentage is variable and it crosses 1, as expected. But even among those patients the two estrogen signatures have a HR below 1 (0.26 and 0.48) and a confidence interval that is not crossing 1. This further shows that ER IHC does not provide a complete description of endocrine therapy sensitivity and scores can replace it at those cases.

## Single sample integration preserves relevant breast cancer properties

Given that not all patients with high ER IHC percentage will respond the same to endocrine therapy, we aimed to develop a framework where we can molecular differences of patients that have similar bulk RNA-seq profile of the tumor tissue. This analysis might unravel changes in molecular pathways that might be used when deciding the treatment. We developed a single sample batch effect removal method (see methods section) to integrate microarray and bulk RNA-seq and create a molecular landscape, which is an embedding of the molecular data into a common space for all samples. The advantage of the method is that given a new sample, it can easily be integrated with all the other previous samples without retraining.

The first step is to normalize the samples to the same scale. The average ranking of housekeeping genes distribution is similar among the two RNA-seq datasets, SCAN-B and TCGA and their mean values are higher than the mean of the distribution on METABRIC (Figure S1 (a)). The normalization preserves the distinctions of gene expression levels between ER+ and ER- breast cancer samples, ESR1 and TFF1 being such examples (Figure S1 (b)).

The biplot in [Figure 2](#fig-02) (a) with the third and fourth components from TCGA and METABRIC samples shows that the samples are overlapping across the two cohorts. All samples, including those used for training and validation, are plotted.

In order to check the robustness of the procedure, we redid all the pipeline 10 times with 10 random subsets of patient samples from TCGA and METABRIC, simulating a cross validation process. The PCA embedding is invariant to rotation, translation and reflection ([Figure 2](#fig-02) (b)).

Missing genes are a problem inherent to publicly available datasets. We try to understand the effect of missing genes in the embedding based on their loading values. Ideally if a low number of genes with high loadings are missing, it should not affect the embedding. On the other hand, the more genes missing with high loadings, the more it will impact the embedding, as PCA takes a weighted linear combination of the gene expression values based on the loadings. We removed 200 genes in total with a varying proportion of top loading genes (ranging from 0 to 100% with a 5% step). The number of top loading genes missing from the dataset is key for the embedding ([Figure 2](#fig-02) (c)). The higher the proportion the less precise the embedding is, with the embedding converging towards the origin, i.e., the (0,0) coordinates ([Figure 2](#fig-02) (c)).

The third component corresponds to the separation between ER+ and ER- BC patients in both cohorts ([Figure 2](#fig-02) (d)). A combination of the third and fourth components shows a good distinction among the PAM50 molecular subtypes ([Figure 2](#fig-02) (e)), showing that the embedding captures important molecular information. The fourth component mostly divides the luminal A and luminal B subtypes, whereas the normal-like subtype is spread across the third and fourth component. This also highlights the fact that one cannot interpret the PCA locations globally, rather when comparing samples one should consider only the neighborhood.

As pointed out before, ER status should be considered continuous and not dichotomous, [Figure 2](#fig-02) (f) shows a gradient of the ER signaling score . The higher values are on the far right of the third component, going to negative values as one goes from right to left, i.e., moving from a more ER+ status to the ER-. Other clinical factors, such as tumor stage, node stage, age, NPI and tumor purity show no influence in the embedding (Figure S2).

## Embedding generalizes to a validation cohort

So far only samples from TCGA and METABRIC were used in the training and projection. An external validation cohort is needed. SCANB was used as an external validation cohort. Similar to the previous results, SCANB is also overlapping with METABRIC and TCGA ([Figure 3](#fig-03) (a)). ER+ and ER- BC patients are well separated ([Figure 3](#fig-03) (b)) and the procedure can also distinguish the molecular subtypes ([Figure 3](#fig-03) (c)) on top of the other samples coming from TCGA and METABRIC. We show that the embedding works for the SMC cohort (34) (Figure S3). As an RNA-seq cohort, it is expected that SCANB samples will be closer to TCGA than to METABRIC when removing batch effects, mostly due to platform. Biplot of PC1 and PC2 ([Figure 3](#fig-03) (d)) shows that SCANB is closer to TCGA than to METABRIC.

The molecular landscape was also validated using patient derived xenografts (PDX) (23). These PDXs are breast cancer cells derived from patient tumor samples obtained from clinics and injected into the mammary gland of mice (35). By using the MIND model, we can engraft cells coming from ER+ BC tumors. Moreover, to have a succesfull engraftment rate, the cells need to grow so they can establish across the ducts. Therefore, the cells when extracted for RNA-sequencing are usually in a proliferative state. [Figure 3](#fig-03) (e) shows the embedding of six different PDX samples with multiple biological replicates (23). These samples are in the scattered around the luminal B region, showing that the landscape captures the biology of the experiment and also it shows the intertumor heterogeneity among ER+ BC patients in a research environment.

## Molecular landscape is a tool to understand and explore patient heterogeneity

Since the molecular landscape relies in a single sample to obtain the embedding, we can add samples from any cohort. The POETIC trial (36) was a trial that evaluated the use of perioperative aromatase inhibitors in ER+, postmenopausal BC patients. Its primary endpoint was time to recurrence. Matched samples from baseline (before treatment) and at surgery (after an average 14 days of treatment) (22) were sent for microarray hybridization. There are also untreated patients, used to control for sample processing artefacts. The patients have matched Ki67 percentage levels, which can be considered an indication of how well a patient responded to the endocrine therapy. Patients with more than 5% of baseline Ki67 and a reduction of 60% upon endocrine therapy are considered responders, otherwise they are called non responders.

In order to gain insights on the differences between responders and non responders, we embedded the POETIC trial samples using the procedure and studied molecular landscape ([Figure 4](#fig-04) (a) left). The samples are spread across the whole molecular landscape, consistent with patients having different molecular biological properties. Furthermore, the POETIC samples are embedded closer to the METABRIC samples (supplementary figure 4; 4.3 from the book). Given the available information, the BC patients that are ER+ and in the left part of the landscape (ER- patients), are all non responders ([Figure 4](#fig-04) (a) right). This highlights the importance to look more carefully to ER+ patients. We selected two patients, a responder and non responder that are close in the embedding ([Figure 4](#fig-04) (b)) to highlight their molecular differences and see what is their context. [Figure 4](#fig-04) (c) shows the average posterior distribution of the neighborhood for the responder patient. The responder patient has a ER signaling score higher than the average. On the other hand, the non responder has a smaller ER signaling score than the average ([Figure 4](#fig-04) (d)) and also a higher androgen signaling score (Androgen response). Other pathways, such as EMT, E2F targets, P53 and TGF signaling along with their average posterior distributions are shown for both patients.

## Generating hypothesis: subgroups and alternative treatments

One key aspect of drug discovery is finding the right groups for targeting with the proper drug. With the molecular landscape we can try to identify different subgroups, based on molecular pathway scores. We identify several pathways driving the distinction between samples in the embedding, such as G2M checkpoint, epithelial to mesenchymal transition, DNA repair and PI3K AKT MTOR signaling ([Figure 5](#fig-05) (a)). The scores go from one direction to other across the two different principal components, indicating that they complement each other when capturing information. Not only G2M checkpoint is differentiating luminal A and B patients, but also EMT.

Understanding a pathway and its clinical relevance is key for drug development and subtyping. The overall and recurrence free survival calculated from patients that have ER+ BC, received only endocrine therapy (METABRIC: OS/RFS and SCANB: OS/RFS) and have show that G2M checkpoint has a hazard ratio higher than 1 ([Figure 5](#fig-05) (b)) and a tight confidence interval, which reflects the fact the higher the proliferation the worse the outcome, and reflects the subtyping differences between luminal A and B. The PI3K AKT MTOR signaling has a hazard ratio higher than 1 with a wider confidence interval, indicating a possible subgroup among all selected patients that could benefit from additional adjuvant therapy targeting this specific pathway. Current clinical trials (37) are evaluating the use of PI3K-AKT-MTOR inhibitors in advanced metastatic BC patients, here we show weak evidence on the molecular level that such treatment could benefit early stage primary BC patients.

# Discussion

Personalized medicine is a key topic in medicine and BC (23). The goal of better understanding the molecular underpinnings of the diseases leads to a better allocation of treatments and resources in the patient care. This has been shown to be necessary by using PDX mouse models, where different PDXs respond differently to hormone treatments (23). We have shown here a possible framework to deal with personalized medicine in breast cancer in general with a focus on ER+ BC patients.

Estrogen receptor status is defined as a clinical variable that usually have two or three categories (3). Breast cancers are classified either in ER+ or ER- based on their protein expression levels and IHC. ER+ are those tumors that express more than 1% and for those expressing less than 1% they are considered ER-. This definition depends on the country and guidelines that are used. For example, the ASCO guidelines (3) recommends to use a threshold of 1%, whereas the Swedish national guidelines uses a threshold of 10% for ER status (38). For those tumors that are ER+, they can be subdivided into low ER positive (1% < IHC% < 10%) and simply ER positive. The low ER positive tumors do not usually benefit as much on endocrine therapy. Here we show using cox regression and multiple big study cohorts (12,20,21) with both RNA-seq and microarray data, that ER status is more of a continuous rather than a categorical state. This can aid when evaluating treatment options and avoiding overtreatment. Several gene signatures already are being used, such as OncotypeDX, Mammaprint and Prosigna to assign chemotherapy for those patients with higher risk of recurrence (5–7). This score might be associated with the commercial signatures, as it has been shown that OncotypeDX’s estrogen module is highly correlated with the general OncotypeDX signature (11).

Integrating molecular data stemming from different sources is a challenge. On one hand batch effect tools are usually able to remove the batch effects across the different sources of variability (17,18), on the other hand they are not single sample based, meaning each time a new sample comes the algorithm needs to be run again. It is also based on the fact one has enough data in the different datasets, otherwise it skews the possible integration towards one of the datasets. Here we show by using TCGA, METABRIC and SCANB that it is possible to integrate the samples from these cohorts in a single sample manner. The samples show good mixing when using test samples not seen during the training stage. The embeddings preserve key molecular features of breast cancer. PC3 is clearly driven by estrogen receptor signaling where from right to left there is a gradient in the ER scores, such as Estrogen early and . On the other hand, PC4 is what makes a difference between the molecular subtypes luminal A and B, which in practice differ by proliferation status in terms of Ki67 levels (39).

Moreover, the embeddings in the validation set (SCANB) preserve the key features of breast cancer. Samples are projected by their different PAM50 molecular subtypes and there is a gradient of estrogen signaling pathway from ER+ BC towards ER- BC patients in all the three cohorts combined. The first two components are the batch effect removal components and SCANB is projected closer to TCGA, since both datasets are RNA-seq. On the other hand, part of the POETIC trial was sequenced using microarray dataset (22,36), and we show that it is projected closer to METABRIC as expected according to the first two components. This highlights that the method can capture information from different technologies and remove such batch effects. Moreover all the samples were sent for sequencing in different contexts at different times and populations, showing the power of the embedding method in removing batch effects and capturing truly the biology of breast cancer.

Sometimes when dealing with publicly available datasets, not all of the genes are available due to ethical protocols (12), pre-processing or technical reasons. Therefore we showed how robust the projection is to missing genes with high loadings in the projection. If less than 20% of the genes we can recover almost surely the position of the embedding if all the genes are available. The more genes that are missing, the closer the projection will be to the origin, i.e., the (0,0) coordinate in the embedding of the third and fourth components of the PCA.

To show the clinical validity of the projection, we used a subset of patients from the POETIC trial with microarray and clinical information (22). The samples are projected as expected and surprisingly the samples that are considered to be non responders upon 2 weeks of aromatase inhibition are projected among the ER- BC patients. When looking further upon two different patients that have similar embedding but different response to endocrine therapy, we see that the responder has a higher value of estrogen signaling than the average. On the other hand, the non responder patient has a smaller estrogen signaling score than the average, which suggests a possible explanation for the difference in response. Moreover, estrogen and androgen receptor signaling have been show to be tightly linked (40) and these two patients have different androgen signaling scores. The non responder has a higher score than the average compared to the responder, whose score is just the same as the average.

Complementing the personalized approach, the molecular landscape can be used as a starting point to understand breast cancer molecular subgroups in a more intuititive way using pathways. Using survival analysis we show that some of the pathways that are important for the embedding can be used for subgrouping and hypothesis generation.

Some weaknesses of the proposed method is that we rely on GSVA scores, which can be used and compared across different cohorts since they have a representation of all molecular subtypes. A way to circumvent this is by having a small library of RNA-seq samples that are representative of the patient population. This way when scoring a new patient, the scores can be compared across different cohorts. There is still a cost barrier for using RNA-seq dataset in the clinical setting, but efforts are being made across the industry to reduce the sequencing costs and make it more widespread. An example is Alithea genomics, a company aiming to provide large-scale RNA-sequencing by using Bulk RNA barcoding and sequencing (BRB-seq).

In conclusion we provide some evidence that ER status should be considered a continuous marker rather than a categorical one. We also extend this notion to a framework for personalized medicine, where each patient is embedded in some context that can be used for the interpretation of its molecular underpinnings. In this paper we only discussed some pathways, but when analysing patient data, several pathways can be considered and should be taken into account when deciding tumor treatment.

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| Figure 1: Scores and survival analysis results from TCGA, SCANB and METABRIC cohorts. (a) GSVA scores for the SET ER/PR signature for each cohort. Each point corresponds to a patient sample and they are divided by estrogen receptor status. (b) Forest plot of the survival analysis for each cohort separately. NPI: Nottingham prognostic index. Ti: i-th stage of tumor. Ni: i lymph nodes with breast cancer cells. |

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| Figure 2: (a) Biplot using the third and fourth components on TCGA and METABRIC samples. Colored by cohort. (b) Same as **a**, colored by ER status. (c) Same as **a**, colored by PAM50 molecular subtype. (d) Hex grid calculated on the biplot of the fourth and third component. Each hex is colored based on its average value of the SET ER/PR signature. (e) Embedding of random samples given different training sets for PCA. Blue dots correspond to the original embedding of a sample and red dots correspond to the new embedding given the new training set. (f) Biplot of all possible embeddings of sample given a certain proportion of top loadings missing in the dataset. |

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| Figure 3: Validation of the molecular landscape with an external cohort (a) Biplot using the third and fourth components and now including all samples from the three cohorts: TCGA, METABRIC and SCANB. (b) Same as **a**, colored by ER status. (c) Same as **a**, colored by PAM50 molecular subtype. (d) Biplot using the first and second component of TCGA, METABRIC and SCANB. |

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| Figure 4: Embedding of the POETIC cohort into the molecular landscape and pathway analysis for patient samples. (a) Biplots of the POETIC samples (baseline and surgery) into the molecular landscape. Left plot is colored by ER status and right plot is colored by molecular subtype PAM50 when available. (b) Biplot highlighting two patients with similar embedding and different response status. (c) Posterior distributions of the average scores in the neighborhood of the responder patient. Red line corresponds to the patient score. (d) Posterior distributions of the average scores in the neighborhood of the **non**-responder patient. Red line corresponds to the patient score. |

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| Figure 5: Average scores in hex regions using all three cohorts together and survival analysis in each cohort individually. (a) Biplots of all samples from TCGA, SCANB and METABRIC that were grouped in small hex regions. Colors are depicted as the average score value in each hex region. Selected pathways are shown. (b) Survival analysis results obtained for each pathway individually. The pathways were adjusted as described in the methods section. CI: confidence interval. |