Integration of Clinical and Gene Expression Data Has a Synergetic Effect on Predicting Breast Cancer Outcome

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Abstract
Breast cancer outcome can be predicted using models derived from gene expression data or clinical data. Only a few studies have created a single prediction model using both gene expression and clinical data. These studies often remain inconclusive regarding an obtained improvement in prediction performance. We rigorously compare three different integration strategies (early, intermediate, and late integration) as well as classifiers employing no integration (only one data type) using five classifiers of varying complexity. We perform our analysis on a set of 295 breast cancer samples, for which gene expression data and an extensive set of clinical parameters are available as well as four breast cancer datasets containing 521 samples that we used as independent validation.

Introduction
Many predictors of breast cancer outcome have been published. These predictors have been derived from gene expression data, such as the 70-gene (Veer et al. [1]), and 76-gene (Wang et al. [2]) signatures, or clinical data, such as the Nottingham Prognostic Index (NPI [3]) and AdjuvantOnline! tools [4]. A few studies have aimed at training a model using both of these data types. In doing so, several approaches were followed, that we outline below.

First of all, the clinical data can be used as a means to stratify patients in subgroups, and then train a gene expression predictor in each of the subgroups. For instance, Wang et al. [2] and Teschendorff et al. [5] have trained a gene expression classifier for ER positive, and separately for ER negative patients [6]. Alternatively, multiple clinical parameters can be used as the initial stratification. For example, Dai et al. [7] stratified into ER/Age-high, and ER/Age-low. Stratifications for ER and HER2 have also been made using gene expression data rather than clinical data, which could lead to better prognostic value [8]. Most of these studies have employed a set of standard clinical variables, such as ER status, tumor grade, tumor size, etc. Horlings et al. [In preparation, [9]] have characterized additional clinical features (e.g., matrix formation, central fibrosis, etc.) for an existing cohort of 295 breast cancer samples [10]. By themselves, these additional clinical variables have independent prognostic power. However, if and how this power can be used to build a better classifier for outcome prediction has not been investigated.

Gevaert et al. [11] have used a Bayesian framework to combine expression and clinical data. They found that decision integration (combination of the outputs of Bayesian classifiers trained on either data type), and partial integration (structure learned per data type, parameters learned after combining the data types) lead to a better performance, whereas full integration (concatenation of the two data types, followed by training the model on the complete set) showed no improvement. These results were obtained by using a cross validation approach on the 78 samples in the Veer et al. [1] dataset. However, on the 19 sample validation set from the same study the pure gene expression based classifier (i.e. no integration) performs slightly better. A major concern in their analysis is that a supervised preselection of genes is performed on the entire dataset, resulting in a potential bias [12]. On the same dataset, Boulesteix et al. [13] employed a random forests and partial least squares
group in the original nominal clinical variable. This way, we obtained a total of \( p_c = 54 \) clinical features. Throughout this paper, we’ll refer to the clinical data as ‘C’.

We applied mean-variance normalization per feature, per dataset (i.e. for both E and C) to ensure approximately equal spread for all features.

### Other Datasets

Reyal et al. [17] have compiled a collection of six datasets, leading to a total of 947 breast cancer samples. From this compendium we have extracted the samples for which Age, Tumor Size, Grade, ER status, Lymph Node status as well as the poor/good survival label (using the same 3 year threshold as for the Vijver dataset) were available. This lead to a total of \( N_e = 521 \) samples (107 poor, 414 good) from the Desmedt et al. [18], Miller et al. [19], Loi et al. [20], and Chin et al. [21] datasets. The NPI was calculated using these clinical parameters as previously defined [22], and both the continuous as well as discretized NPI were appended to the clinical data. Thus, a total of seven clinical parameters, \( p_c = 7 \) (this is much less than the \( p_c = 54 \) in the Vijver dataset), were available for all 521 samples. For the expression data we used the probes that were also present in the Vijver dataset, by matching Entrez ids (\( p_c = 11601 \)). After this selection, we applied mean-variance normalization per feature, per dataset (i.e. for both E and C).

### Classifiers

We employed five classifiers with varying degrees of complexity, some of which have been used before to integrate clinical and expression data. We shortly discuss each classifier (see Table 1):

1. **A Nearest Mean Classifier (NMC), with the cosine correlation as distance measure.** This linear classifier has previously been applied on expression data, and was shown to outperform more complex classifiers [1,23].

2. **A Simple Bayes Classifier (SBC) [24], which is based on the assumption that the features are independent.** This simplifies the computation of the class conditional densities significantly. In spite of this simplification, it has been shown that this classifier performs remarkably well [24]. Class continuous densities of continuous features were modeled using Gaussian distributions.

### Materials and Methods

#### Vijver Dataset

We have used the 295 breast cancer sample dataset from Vijver et al. [10]. For all \( N_e = 295 \) samples microarray data is available. We selected the \( p_e = 15676 \) probes with an Entrez identifier. From this dataset, we selected \( N_e = 259 \) samples, which we could assign to a poor/good outcome group based on their survival characteristics (poor: event within five years of follow up, good: at least five years of metastasis free survival), a dichotomisation commonly made, e.g. Veer et al. [1]. Thus, the remaining 36 samples were not included in the dataset since these have been censored before five years of follow up, making it impossible to assign them to the correct outcome group. Throughout this paper, we will refer to the expression data as ‘E’.

In addition to expression data, we have a variety of clinical data available (Horlings et al., In preparation, [9]). The clinical features include the originally published variables (e.g. Grade, Age, ER status, etc), outputs from clinical models (e.g. NPI, StGallen, and Adjuvant), complemented with a set of novel pathological variables (e.g. Matrix Formation, Central Fibrosis, etc.). Table S1 shows a complete list and details of the clinical variables used. In total, we considered 45 clinical variables (which have no missing values for these \( N_e = 259 \) samples), of which 2 were nominal, 33 were binary or ordinal, and 10 were continuous. The two nominal variables were converted into binary features, i.e. one feature per
3. A 3-Nearest-Neighbor classifier (3NN) [25], employing the cosine correlation as distance measure, and majority voting to assign a sample to a class. Since there is a class imbalance, the majority vote is adjusted with the class priors. This classifier is capable of constructing non-linear decision boundaries. Moreover, it is frequently applied to microarray data, see e.g. Dudouit et al. [26].

4. A Support Vector Machine (SVM) [27], using a cosine correlation kernel [14], i.e. a kernel function which computes the cosine correlation between two input objects. This classifier is appropriate for small sample size problems, and has previously been used to integrate expression and clinical data [14]. The cosine correlation kernel for SVMs is identical to a linear kernel, where the feature vector for each sample has been divided by its L2-norm [28]. The C parameter was fixed at 1 (default value, svmtrain in Matlab R2012a). To account for class imbalance, C was rescaled by \( N_c / (2N_{poor}) \) for the samples in the poor group and by \( N_c / (2N_{good}) \) for the samples in the good group.

5. A Tree classifier (Tree) [29], which allows for highly non-linear decision boundaries. Gini’s diversity index was used as splitting criterion. In order to regularize the tree classifier, we employed two variants. The first variant (Tree1), optimizes the tree depth but selects a subset of features from all features. The second variant (Tree2) is not pruned, but selects features from the subset of up to 200 most predictive features as provided by the feature selection procedure.

We excluded the Bayesian approach introduced by Gevaert et al. [11], since it is computationally intractable to train this model on all genes.

Cross Validation Setup

To evaluate the performance of the classifiers, and determine the optimal number of features (tree depth for the Tree and the HybridTree classifiers (see Section ‘Integration strategies’)), we applied a double loop cross validation protocol (DLCV, Wessels et al. [23]). The DLCV procedure employs two loops, an outer loop for validation purposes to estimate the performance on a left, out independent part of the data, and an inner loop in which the classifier’s parameters are optimized. The DLCV procedure can be described in a few steps:

1. For each repeat, the data is split (stratified) into five parts (different splits for each repeat).
2. For each fold, four parts are used for the inner loop (training set), the fifth part is used in the outer loop for validation (validation set).
3. On the training set, a 10-fold cross validation is performed to estimate the optimal number of features \( n \) is defined as the number of features at which the \( e_{FPFN} \) is minimal) to be used in the classifier, i.e. the number of features that resulted in the best classification performance based on the 10-fold cross validation.
4. Next, a classifier is trained on the complete training set, using the estimated optimal number of features.
5. Finally, the performance of that classifier is assessed on the validation set.

Typically, datasets are imbalanced in the sense that the samples from the classes do not appear in equal fractions in the dataset. Moreover, the imbalance will be different for different datasets.

Hence, directly comparing overall error rates (fraction of wrong assignments), is not an appropriate comparative measure. Therefore, classification errors were calculated by using the average False Positive False Negative ratio, defined as:

\[
e_{FPFN} = \frac{FN + FP}{2}
\]

where \( TP \) represents the number of true positives, \( TN \) the number of true negatives, \( FP \) the number of false positives, and \( FN \) the number of false negatives. This ratio is equivalent to 1 - \( \frac{TP}{TP + FP} \) (Sensitivity + Specificity).

The entire protocol was run 60 times (i.e. 60 repeats of the double loop cross-validation protocol). To find the optimal number of features, we constructed learning curves in the inner loop for up to 200 features (or 54 when only using the clinical data). In all experiments, we used the exact same repeats and folds.

As a result, we were able to compare the performance results in the outer loop on a pair-wise basis, using a one-sided, paired t-test.

Kaplan-Meier curves were constructed by using the predictions that were made in the outer loop. Consequently, in each repeat, every sample has once been part of the test set in the outer loop. Thus, for each sample we have a fully unbiased prediction of the binary label. After completing the 60 repeats, we have 60 unbiased predictions of each sample. Next, we take the mean of those 60 predictions, and assign a sample to the poor group if the average is below .5 and to the good group when the average is above .5. This approach is known as the ‘pre-validation strategy’ [30]. The predictions are independent, but nevertheless the training sets will overlap in terms of samples. However, this only yields a small bias [30].

As an alternative performance criterion, we also considered the AUC (Area Under the Curve) of the ROC (Receiver Operator Characteristic) curve instead of \( e_{FPFN} \). We employed the perfcurve function in Matlab, which tests all possible thresholds on the vector of classifier output scores, and then uses trapezoidal approximation to estimate the area under the curve (AUC). The ROC analysis can straightforwardly be applied to the ‘early’, ‘intermediate’, and ‘no integration’ setups. Using the vector of scores obtained from the classifier, we varied the threshold in steps of 1 sample. However, the late integration setups require two binary vectors, and thus require choosing an operating point on each of the two separate classifiers. This complicates the construction of an ROC curve. We solved this problem as follows. Each classifier outputs a ranking of the samples from most likely to least likely poor outcome. For \( N \) samples this results in a total of \( N^2 \) possible thresholds (ROC operating points) for the joint classifier. Rather than considering all these possibilities, we only considered operating points where both classifiers assign the same number of samples to the poor (and good) outcome class, resulting in \( N \) joint operating points. So, for the \( i \)-th operating point, we set the threshold on both the the E and C classifier such that \( i \) samples are classified as poor outcome. This results in two binary vectors, both with \( i \) values set to 1 [poor outcome] and the rest to 0 [good outcome]. After that the two vectors of binary prediction labels are combined using the AND/OR operator, and compared against the true label to provide the sensitivity/specificity coordinates for the ROC curve.

Feature Selection

In the inner loop of the cross validation procedure, we used a feature filtering approach. To rank the features, we employed a t-test for the continuous features and the chi-squared test for discrete features. The combined set of features are then ranked based on the p-values of the associated tests.
Integration Strategies

Following Gevaert et al. [11], we considered early, intermediate and late integration. In Figure 1 we depict each of the three strategies, and describe them below. Table 1 shows which integration strategies are considered in combination with which classifiers.

Classifiers are indicated by their abbreviation, followed by the type of integration used. For example, ‘NMC; None E’ is the nearest mean classifier (NMC) without integration (None), trained on expression data (E).

Early Integration

For the early integration strategy we concatenated the E and C datasets, and thereby created a single dataset, EC, with \( p_e + p_c = 15730 \) features. Classifiers trained on EC are indicated with the suffix ‘Early’, e.g. ‘NMC; Early’ for the NMC variant.

Intermediate Integration

For all classifiers (except the intermediate Tree classifier), we determine the optimal sets of features from each data type separately (in the inner loop). In all subsequent steps, we used these sets of features and all training samples. We define \( d \) as a mixing parameter (ranging from 0 to 1) and \( d(a; b; c) \) as the cosine correlation between vectors \( a \) and \( b \) using the optimal features in \( c \). What we do for each classifier is described below:

1. For the NMC classifier, we compute the centroids of the poor and good class (denoted as \( c_{poor} \) and \( c_{good} \)) for both the E and C data types separately. Next, for a sample \( x \), we compute a combined distance \( d_{NMC} \) to the centroids, which is a linear function of the distances in the individual spaces, and is formulated as:

\[
d_{NMC}(x, c_{poor}; E, C) = \sum d(x, c_{poor}; E) + (1 - \alpha) d(x, c_{poor}; C),
\]

2. For the SBC classifier, we first calculated the distance of a sample \( x \) to a training sample \( y \) in E and C, leading to \( d(x, y; E) \) and \( d(x, y; C) \). Next, the overall distance \( d_{SBC} \) is computed as a linear combination of the individual distances:

\[
P_{SBC}(poor|x; E, C) = \alpha P_{good|x; E} + (1 - \alpha) P_{poor|x; C},
\]

3. For the 3NN classifier, we first calculated the distance of a sample \( x \) to a training sample \( y \) in E and C, leading to \( d(x, y; E) \) and \( d(x, y; C) \). Next, the overall distance \( d_{3NN} \) is computed as a linear combination of the individual distances:

\[
P_{SBC}(good|x; E, C) = \alpha P_{good|x; E} + (1 - \alpha) P_{poor|x; C}.
\]

Subsequently, the sample \( x \) is assigned to the class for which the distance \( d_{SBC} \) is the smallest.

Figure 1. Schematic indication of the expression dataset (E), clinical dataset (C), along with different integration strategies that were tested. Examples are shown for the NMC classifier. On the left, we depict the ‘no integration’ setup, for which a separate classifier is trained on each dataset (‘NMC; None E’ and ‘NMC; None C’). For early integration, the two datasets are concatenated into EC, on which a single classifier is trained (‘NMC; Early’). Similarly, for intermediate integration, the datasets are combined at an intermediate step in learning the classifier (‘NMC; Intermediate’). Finally, late integration is depicted on the right, where a classifier is trained on each dataset separately, and combined by means of a logical function (‘NMC; Late OR’). doi:10.1371/journal.pone.0040358.g001
three closest samples (the majority vote is adjusted with the class priors).

4. For the SVM classifier, we use the cosine correlation to compute a kernel $k(x,y;D)$, the kernel distance between samples $x$ and $y$ given data type $D$. We then construct a new kernel matrix $k_{SVM}$ by taking a linear combination of the kernel matrices from the separate data types:

$$k_{SVM}(x,y;E,C) = ak(x,y;E) + (1-a)k(x,y;C).$$  \hspace{1cm} (7)

For computational tractability, positive semi-definiteness of the kernel has to be ensured (Mercer conditions). This is the case when the weights employed in the linear combination (7) are non-negative [14], a condition which is satisfied here (cosine correlation as kernel).

5. For the Tree classifier, we followed an approach similar to Pittman et al. [16]. First, we considered a method where we start with a NMC trained on C (since this is a computationally inexpensive classifier with known good performance). This classifier splits the samples into two groups, each associated with a node in the tree. In these and all subsequent nodes, we branch further using a NMC trained on E and the samples at the relevant node. The procedure was stopped when a particular branch was pure (only poor or only good samples), or contained fewer than ten samples. This approach will be referred to as HybridTree (C). We also included the complementary setup, which starts with a NMC trained on E, and uses NMC classifiers trained on C in the subsequent nodes (HybridTree (E)).

For the two HybridTree variants, we optimized the tree depth in the cross validation procedure (inner loop), while we fixed the number of features used in each classifier to the top 100 features when trained on E, and the top ten features when trained on C.
These features were selected using all training samples (in the inner cross validation loop) in a particular branch of the tree, using the same feature selection methods as described above in the Section 'Feature selection'. The mixing parameter $\alpha$ that several intermediate integration strategies use, is also optimized in the inner loop, now using the entire training set. More specifically, we vary $\alpha$ from 0 to 1 in steps of 0.01, and then inspect the error on the training data. This ensures that the $\alpha$ parameter is optimized in an unbiased fashion since the test samples in the outer loop are not involved in optimizing $\alpha$.

Classifiers trained using the intermediate integration strategy are indicated with the suffix 'Intermediate', e.g. 'NMC; Intermediate' for the NMC variant.

**Late Integration**

For late integration we train a classifier on E and C separately. After that, we apply a logical function on the binary classifier outputs (poor is positive, and good is negative).

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**Figure 3. Overview of all pairwise comparisons of the classifiers.** Comparisons were made by means of a one sided, paired t-test, testing the hypothesis that the error associated with the approach listed in the row is lower than the error associated with the approach listed in the column. Red cell shading indicates a p-value smaller than 0.05, and white cell shading indicates that the p-value was larger than 0.05. Letters in the cell refer to particular comparisons that are discussed in the text. 
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considered a logical AND function, for example for the NMC classifier:

\[ \text{NMC; Late AND} = 'NMC; None E' \ AND \ 'NMC; None C', \]  
(8)

and a logical OR function, for example for the NMC classifier:

\[ \text{NMC; Late OR} = 'NMC; None E' \ OR \ 'NMC; None C'. \]  
(9)

The difference between the logical AND and OR functions is the way the discordantly classified samples are treated. Using the AND function these are assigned to the good class, and using the OR function these are assigned to the poor class. These two options are formally known as ‘believe the positive’ (OR) and ‘believe the negative’ (AND) integration [31].

**Results**

The ‘NMC; Late OR’ Classifier Performs the Best

The lowest error is achieved using the NMC classifier with late OR integration strategy (‘NMC; Late OR’, eFPFN = 0.273, Figure 2A. Figure 3 (squares indicated with an ‘A’) shows that this error is significantly lower than all other classifiers. This is a clear indication that there is synergy between the two data types, and that the late OR integration strategy provides a way to exploit the synergy. In addition, Figure 2B shows that the Kaplan-Meier curve of the ‘NMC; Late OR’ classifier is more significant than those from the NMC classifiers trained on a single data type. More specifically, the good group has become purer at the five year point (94.4% metastasis event free, versus 87.5% and 88.6%, respectively).

Figure 4 shows the Kaplan-Meier curves of four other signatures that were applied to the same set of 259 samples from the Vijver dataset (70-gene signature, Veer et al. [1]; 253-gene hypoxia signature, Chi et al. [32]; 186-gene invasiveness signature, Liu et al. [33]; 97-gene genomic grade index signature, Sotiriou et al. [34]). The p-value of the ‘NMC; Late OR’ Kaplan-Meier curve is lower than each of these other four signatures. That is, the ‘NMC; Late OR’ strategy performs comparable to or better than all these

![Figure 4. Kaplan-Meier curves of the same 259 sample subset from the Vijver dataset, employing four different signatures. P-values reflect the logrank test. doi:10.1371/journal.pone.0040358.g004](image4.png)

![Figure 5. Boxplot showing the α values that are obtained using the different classifiers with an intermediate integration strategy (300 α values from the 60 repeats of 5 folds). doi:10.1371/journal.pone.0040358.g005](image5.png)
signatures as measured by either the significance of the log-rank p-value or the fraction of patients that remain metastasis free at 10 years. This is especially noteworthy in the case of the 70 genes as this signature was trained on a subset of de Vijver dataset, and is therefore expected to be positively biased.

Integration Improves Performance

The NMC, SBC, SVM, and Tree1 classifiers perform the best when employing the late OR integration strategy, whereas the 3NN classifier performs the best when employing the early integration (see Figure 3, squares indicated with a ‘B’). In addition, the median mixing parameter $\alpha$ that was selected in the intermediate approaches is around .5 or higher (see Figure 5), suggesting that both data types are important. Thus, integration of the two data types proves beneficial for all classifiers with the ‘Late-OR’ strategy resulting in the best performance for all classifiers except the 3-NN classifier.

Less Complex Classifiers Outperform Complex Classifiers

Figure 3 shows that the NMC classifier outperforms all other classifiers, with the exception of the ‘SBC; Late OR’ option. Overall, we can approximately rank the classifiers based on the achieved error rates in the following order: NMC < HybridTree < SBC < 3NN < SVM (with the remark that the C parameter of the SVM has not been optimized) < Tree. This ordering correlates with the complexity of the classifiers, and confirms previous results [23,35]. The most likely explanation for this ordering is the small sample size problem, due to which the more complex classifiers run into overtraining problems, and consequently perform worse on independent data.

A Hybrid Tree Approach is not Useful on Breast Cancer Datasets

The average tree depth that is selected when using the HybridTree (C) classifier is 1.1. At this tree depth, the HybridTree (C) is practically equivalent to the NMC using clinical data. On the other hand, the HybridTree (E) has an optimal tree depth of 1.8. This suggests that a second level of NMCs using clinical features might be beneficial on top of the expression NMC. However, both HybridTree classifiers are significantly outperformed by the NMC classifiers without any integration (Figure 3, indicated with a ‘C’). We suspect that this is due to the extremely small numbers of samples available in the second layer and further down the tree. The classifiers in these nodes are most likely highly overtrained and consequently do not generalize very well.

The HybridTree (C) setup is very similar to training an expression based classifier within clinical subgroups. Our analysis indicates that, there is little to be gained by such a strategy. The intermediate and late integration strategies using a NMC are better options.

Expression and Clinical Features Perform Equally Well

A major selling point of existing gene expression based classifiers is their superior performance compared to the existing clinical models. However, we observe a small performance advantage for the NMC trained on C compared with the NMC trained on E (Figure 2). This difference isn’t significant, see Figure 3 indicated with a ‘D’. We claim that this might be explained by the more extensive set of clinical parameters that we used. To test this, we split the clinical features into three groups (see Table 1): Original (O, those available at the time the first signatures were published, e.g. grade, age, ER status, etc.), Signatures (S, outputs of clinical
models, e.g. NPI, StGallen, etc.), and New (N, those not published before, e.g. matrix formation, central fibrosis, etc.). We repeated the classification experiments, using an NMC with these different sets of clinical variables. We already tested an NMC using the O+S+N features (‘NMC; None C’), and added NMCs using the O+S features, O+N features and only the O features. The result is shown in Figure 6A. Indeed, the NMC classifier using only the original features (O), performs significantly worse than all other options (Figure 6B). Adding the outputs from the clinical models or new features improves the performance (variants using O+S or O+N), and using all three (O+S+N) gives another large improvement. Thus, by including the outputs from clinical models and the new set of clinical features, the performance of the NMC trained on all clinical features is equivalent to that of the NMC trained on E. Therefore, there is no significant performance argument to choose one over the other.

The Selected Features

The NMC with late OR strategy performs the best, and therefore we trained a final classifier on all samples of the Vijver dataset. The number of features was chosen by averaging the number of features that was found to be optimal in the inner loops, resulting in 87 and 9, for the expression and clinical features, respectively (see Figure 7 for a pairwise correlation of all features).

First of all, we performed an enrichment analysis for the 87-gene signature. We collected gene sets from GO, KEGG, Reactome, WikiPathways, and the Molecular Signature Database C2 (MSigDB), giving a total of 4525 gene sets with at least five genes. We used the hypergeometric test to assess the significance of the overlap, followed by a Bonferroni correction. A heatmap of the enrichment is shown in Figure 8. The most highly enriched gene set is the van’t Veer signature [1] from MSigDB. This is to be expected, since there is sample overlap between the datasets from Veer et al. [1] and Vijver et al. [10] (nevertheless it is a positive control). Other than that, many proliferation associated gene sets are enriched. This has previously been identified as a category picked up by most signatures [17].

The nine clinical variables that were selected are shown in Table 2. The set of clinical variables includes a proliferation signature (Mitos, Grade02, Grade07). Moreover, it contains some
of the hormone associated variables known to be associated with survival (ER, ERbin, PR). In addition, the outputs from some of the clinical models were selected [NPI Score, Clin NPI]. Matrix has not been previously associated with survival.

In order to see whether the clinical features pick up a signal different from the expression features, we inspected their correlation. Figure 7 shows three Subgroups ‘A’, ‘B’ and ‘C’ of correlated genes. Subgroups A and B are correlated with the grade/signature clinical variables (Mitos, grade02, grade07, NPI-score, Clin-NPI). In addition, the smaller set of genes in Subgroup C, are correlated with the ER and PR clinical variables (ER, ERbin, PR). This Subgroup is anti-correlated with Subgroups A and B. We performed the same enrichment analysis on these three subgroups of genes, see Figure 7. The genes in Subgroup A are clearly highly enriched for proliferation associated gene sets, which also confirms the positive correlation with proliferation associated clinical parameters.

The SCUBE2 gene from the smaller Subgroup C is part of the ‘Estrogen genes’ in the signature from Paik et al. [36]. This explains the positive correlation with the ER and PR clinical parameters. However, the genes in Subgroup C are not enriched for any gene sets (see Figure 7). ER regulates many genes, resulting in very large ER associated gene sets. As a consequence, the set of genes in Subgroup C is probably too small to be able to become significantly enriched.

The matrix variable is not correlated with any of the 87 genes, nor with the other eight clinical variables. Next, we tested whether any of the genes is associated with the matrix variable, by means of a t-test. After Bonferroni correction, none of the genes have a significant p-value (at p < 0.05). Thus, the information of the matrix variable is not captured by the expression data at all.

Integration Also Improves Performance on Four Independent Breast Cancer Datasets

The amount of clinical data that is published for breast cancer microarray datasets is often limited. Therefore, a direct validation of the ‘NMC; Late OR’ classifier on independent data is impossible (due to missing clinical features). However, from a previously gathered collection of breast cancer datasets [17], we extracted a total of 521 cases for which survival and seven clinical variables were present (see Materials and Methods section). The NMC classifier with all integration strategies was applied on this dataset, employing the DLCV procedure with the same settings as used for the Vijver dataset (see Materials and Methods section). The other classifiers were omitted since the NMC classifier performed best on the Vijver dataset.

Figure 9a shows the DLCV error rates, and Figure 9b shows their pairwise comparison, revealing that the ‘NMC; Intermediate’ strategy performs the best, followed by the ‘NMC; Late OR’ and ‘NMC; Early’ strategies. Thus, the integration strategies also improve the performance on these four independent datasets. Moreover, the NMC classifiers trained using the expression or clinical data alone perform equally well (eFPFN of 0.342 vs 0.345, no significant difference). Figure 9c shows the Kaplan-Meier curves of these classifiers trained using expression or clinical data alone, showing very similar curves. In addition, employing the ‘NMC; Late OR’ strategy primarily provides a purer good group (0.948 vs 0.903 and 0.894 respectively). The superior performance of the integration strategies, and the equivalent performance of the expression and clinical features confirm our findings on the Vijver dataset.

Integration Results in Higher AUC Performance

In the DLCV procedure, we optimized the number of features by minimizing the eFPFN error. As an alternative, we repeated the experiments aiming to maximize the AUC, which reflects the performance across the entire ROC curve rather than a single operating point (see Materials and Methods). We repeated the experiments using the NMC classifier, as that classifier achieved the best performance in the eFPFN experiments. All DLCV settings were kept the same (60 repeats, 5 folds, etc.). Figure 10 shows the average AUC results, a pairwise comparison of the classifiers, and boxplots of the AUC results. On both the Vijver dataset, and the independent validation datasets, a late integration strategy achieves the highest AUC. Thus, we conclude that integration also improves the AUC performance.

Discussion

For all classifiers tested, we found evidence to support the hypothesis that integration of expression and clinical data leads to better predictors. We hypothesize that this is the result of two effects. First of all, both individual classifiers pick up a noisy proliferation associated signal, and their redundancy leads to a better prediction. Secondly, the clinical set of features has some additional information, for example the ‘Matrix formation’ variable, which is not captured by the expression. This complementarity of features results in a synergistic effect on the classification performance.

The late OR integration is the strategy that most often leads to the best performance improvement on the Vijver dataset. Using the late OR strategy, samples for which the individual classifiers are discordant are assigned to the poor outcome group. As a result, the identified good group becomes smaller but also purer. We hypothesize that this is also why the performance increases, the two data types are primarily synergistic in finding a pure group of good cases. A similar effect was seen when combining the classifier.
outputs of existing gene expression signatures [17]. The intermediate and late OR integration strategy perform the best on the four independent datasets. On these datasets, the late OR strategy also results in a clear improvement in the ten year survival of the good group. Identifying a very pure good outcome group may clinically be the most interesting, since those patients could be spared treatment.

Using the eFPFN as criterion shows that the Late OR strategy is the best on the Vijver data, and the intermediate strategy on the independent data (the Late OR is second best). When using the AUC as criterion, the Late OR strategy performs the best on the Vijver data, and the late AND strategy on the independent data (the Late OR is second best). These differences in the best integration strategy may be due to 1) potential differences in composition of the samples between the cohorts, 2) the use of different microarray platforms, 3) differences in clinical data that is available (much more extensive for the Vijver dataset), and 4) differences in annotation (such as differences in grading between pathologists). Some or all of these effects will play a role in which classifier/integration strategy performs the best. Remarkably, in all cases best performances are achieved by integrating the two data types, showing strong evidence of their synergy.

In the intermediate and late integration strategies, the optimal sets of features are selected on each data type separately and not in the context of the final integrated classifier, which might be suboptimal. We did not explore alternative feature selection procedures, which take this complementarity into account, due to the additional computational complexity.

The nearest mean classifier significantly outperforms all other classifiers. Thus, our results support earlier indications that a relatively simple classifier is least hampered by the small sample size problems. On top of that, we conclude that this is the case regardless of the choice of integration strategy. We would like to stress that these claims can only be made for the breast cancer data sets examined in this study.

Gevaert et al. [11] also investigated the three types of integration strategies, albeit with only one classifier (Bayesian network). Their conclusion that intermediate and late integration perform better are confirmed in this study. In addition, we show that this is the case without preselecting genes, without discretizing the expression data, and on a larger dataset.
Daemen et al. [14] also employed the SVM with intermediate integration, using the same type of kernel (cosine correlation distance). They conclude from their AUC measurements that the SVM trained on clinical data alone performs better than the SVM using intermediate integration, which, in turn, performs better than the SVM trained on the expression data only. Our results show the exact same order in performances. In addition to that, we also conclude that the SVM intermediate and clinical only perform significantly better than the SVM on expression data only. The best option identified in our study, an SVM with late OR integration, was not tested by Daemen et al. [14]. However, our analysis convincingly shows that the choice of using an SVM with this type of kernel is rather poor for this type of dataset, since it is outperformed by several other classifiers.

‘Hormone related’ and ‘Proliferation’ features are selected by both the E and C classifiers indicating the importance of these processes in predicting breast cancer outcome. Matrix formation was selected on the Vijver dataset but was not available on other validation datasets. Scoring additional histo-pathological features on tumor specimens may yield further improvement in breast cancer outcome prediction and is therefore worth pursuing.

Supporting Information
Table S1 Overview and details of the clinical variables used for the Vijver dataset.
(XLSX)

Author Contributions
Conceived and designed the experiments: MHV HMH MJV MJTR LFAW. Performed the experiments: MHV. Analyzed the data: MHV MJTR LFAW. Wrote the paper: MHV MJTR LFAW.

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