Validation of Biomarker-Based Risk Prediction Models
Jeremy M.G. Taylor,1 Donna P. Ankerst,2 and Rebecca R. Andridge1

Abstract
The increasing availability and use of predictive models to facilitate informed decision making highlights the need for careful assessment of the validity of these models. In particular, models involving biomarkers require careful validation for two reasons: issues with overfitting when complex models involve a large number of biomarkers, and interlaboratory variation in assays used to measure biomarkers. In this article, we distinguish between internal and external statistical validation. Internal validation, involving training-testing splits of the available data or cross-validation, is a necessary component of the model building process and can provide valid assessments of model performance. External validation consists of assessing model performance on one or more data sets collected by different investigators from different institutions. External validation is a more rigorous procedure necessary for evaluating whether the predictive model will generalize to populations other than the one on which it was developed. We stress the need for an external data set to be truly external, that is, to play no role in model development and ideally be completely unavailable to the researchers building the model. In addition to reviewing different types of validation, we describe different types and features of predictive models and strategies for model building, as well as measures appropriate for assessing their performance in the context of validation. No single measure can characterize the different components of the prediction, and the use of multiple summary measures is recommended.

Risk prediction tools combining biomarkers with other risk factors are increasingly being proposed for a variety of purposes in the management of individual cancer patients. Many of these tools are available as user-friendly calculators on the internet and easily found and accessed by patients. They facilitate informed decision making between doctor and patient on whether or not to pursue more invasive diagnostic testing, likelihood of progression of disease, or outcome to specific therapies. Some examples include the Prostate Cancer Prevention Trial (PCPT) calculator for predicting the probability of prostate cancer (1), the Gail model, which gives the lifetime risk of breast cancer (2), the many nomograms for predicting disease progression in prostate cancer (3), and Oncotype DX for predicting risk of recurrence in breast cancer based on gene expression data from 21 genes (4). Models that use serial measurements of biomarkers include ROCA for predicting ovarian cancer based on serial measurements of CA125 (5), and a Web-based calculator for predicting prostate cancer recurrence using serial prostate-specific antigen (PSA) values after radiation therapy.3 Combined with the recent explosion of high dimensional panels of markers, including those proposed by genomic, proteomic, and other -omic fields, the uncontrolled proposal and mass posting of risk prediction tools raises deep concerns whether these tools have in fact been suitably validated on external populations of the type intended for use. Indeed there is an “emerging breed of forensic statisticians” (6) debunking the claims of some of these tools; recent analyses of a few high-profile risk prediction tools have shown lack of reproducibility (7, 8). This article educates and/or reminds practitioners of the essential ingredients for proper validation of cancer risk prediction tools, summarized in Table 1. Throughout, we use methods and results from the PCPT and a lung cancer gene expression study predicting survival time from gene expression data (9) to illustrate steps in the validation process.

What is a Model?
In this article, we do not distinguish between prognostic and predictive models (10), terminology that is becoming familiar to oncologists and is described in more detail in the articles by George (11) and Simon (12) in this issue. Prognostic models evaluate general risk, whereas predictive models assess who may respond to a certain therapy. In statistical terms, the distinction is that predictive models include an interaction between biomarker and treatment. Despite this distinction, the general principles behind validation of both types of models remain the same.

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There is vast statistical literature on how to build predictive models (e.g., refs. 13, 14). Multivariable models allowing simultaneous association of biomarkers and predictors with clinical outcome, such as logistic regression for presence/absence of disease or Cox regression with survival, are common building blocks of biomarker-based risk prediction tools. More algorithmic statistical models, such as support vector machines or neural networks, can also be used.

Some models have a simple structure, combining information from a few variables in a transparent way. At the opposite end of the spectrum are complex “black box” models, e.g., a Web-based calculator with a large number of input variables. Some models give a simple prediction such as high versus low risk, more refined models give a continuous score; even more desirable is when the continuous score is on an interpretable scale such as a probability of recurrence. The PCPT calculator uses a multivariable logistic regression model and four variables (PSA, digital rectal examination, history of prostate cancer in a first-degree relative, history of a prior negative biopsy with prostate cancer status on biopsy) to provide an estimated risk of prostate cancer ranging from 0% to 100%. Other examples of simple models are the nomogram models of Kattan et al. (3) that provide an easily interpretable prediction. The graphical approach of the nomogram explicitly shows how three clinical measures (PSA, clinical stage, and Gleason Score) are scored and summed to yield a predicted 5-year cancer free survival probability. An example of a more complex model comes from the lung cancer gene expression study where expression values from 13,838 genes are combined in a weighted linear sum, with the weights for each gene estimated using a penalized Cox model. Prediction is a continuous score, with higher scores indicating increased hazard of death, but the actual score does not have an easy interpretation (i.e., it is not a probability).

However appealing the idea of transparency, simplicity of the model is not necessarily a virtue; performance of the model is more important, and simplicity over complexity should not be the primary consideration in the model building process. Proper validation of a reasonable model is more important than the pursuit of an ideal model, which may never be found in any case. In the lung cancer gene expression study, the chosen model (using 13,838 genes) had slightly better performance in validation data sets than models based on significantly reduced numbers of genes, and the model based on the single best gene did very poorly. However, one benefit to simple models is a logistical one; there may be reduced expense and more focused quality control if the number of biomarkers in the model is small.

In building a model, there is sometimes the perception that newer technologies, such as neural networks, offer a better approach. Although such models are certainly more complex, involving combinations of interactions and nonlinearities, there is also more danger of overfitting and illogical resulting models (15). Whether a model fits better will depend on the context and the available data. In a review of 28 studies directly comparing neural networks to traditional regression modeling, Sargent et al. (16) found that neural networking was not a better approach in more than half the studies. Whatever method is proposed, it is imperative that sufficient details are provided to both understand how it was developed and so that others could implement it themselves.

Models involving biomarkers. Models must be built based on good statistical principles and, to this end, the resulting model should fit the data used to develop it. However, if the predictive model involves a panel of biomarkers then the problem associated with building models with a high number of variables can arise, in which the model overfits the data. Often there is little a priori knowledge about which biomarkers to include in the model, and the temptation is to throw them all into an automated procedure such as stepwise regression. This often leads to the model capturing not only real patterns but also idiosyncratic features of the particular data set, resulting in poor performance in external validation. Better methods for handling data sets with large numbers of variables exist, including penalized regression methods such as ridge regression and LASSO (14). These can be very useful with large numbers of input variables and can provide more reliable results and help avoid overfitting. However, even with more advanced statistical methods for model building validation of a prognostic model is necessary to separate true associations from noise.

In addition to using caution to avoid overfitting, models that include biomarker data can also suffer from low power because of misunderstanding about what drives the power to detect significant effects. Sample size requirements for validating a new biomarker are demanding and even more demanding for showing it to be beneficial above and beyond existing markers. It is not the number of measurements per subject that drives power (e.g., number of genes measured) but the number of subjects. Obtaining tons of measures on a small set of patients will likely not yield significant results, especially for predictive models associated with a specific therapy as defined above. Such models by definition include interaction terms, and thus, large numbers of subjects are required to find any significant interaction effects with treatment.

In our opinion, there is too much emphasis on the specific variables that are included in a prediction model. For example, for Oncotype DX or the lung cancer gene expression study, one may ask, “Why these genes?” or “Why these weights?” In situations where there are a lot of biomarkers to choose from, it will frequently be the case that many subsets of variables give very similar predictions; thus, too much sanctity should not be placed on the specific model. It is not at all surprising in gene expression studies that models built from different data

<table>
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<tr>
<th>Table 1. Components of biomarker-based model validation</th>
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<tr>
<td>- Ascertainment of true outcome</td>
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<td>- Proper model building, including internal validation</td>
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<td>- Avoidance of overfitting in the model building</td>
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<td>- Consideration of biomarker assay reproducibility</td>
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<td>- External validation is crucial</td>
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<td>- External validation necessary for determining generalizability</td>
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<td>- Choosing appropriate performance measures dependent on outcome and prediction type</td>
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<td>- Avoiding overuse of cut-points in risk prediction</td>
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sets for the same goal have contained very different sets of
genes (17).
Models involving biomarkers as input variables raise unique
issues due to technological advances and assay inconsistencies,
discussed in the article by Owzar et al. in this issue (18),
Hammond et al. (19), and the REMARK guidelines (20). Assays
to measure biomarkers evolve over time; thus, measurements
from a new assay cannot be substituted into a model built using
measures from an earlier assay unless the two assays are highly
 correlated. If correlation is weak, then the performance of the
predictive model will likely be worse even if the new assay
provides more accurate measurement. Even when changes in
technology do not interfere, any number of factors may cause
an assay to be systematically higher or lower or more variable in
one lab compared with another.
One caveat in prognostic model building is that small
P values can be misleading. An often disappointing result of
multivariable risk models combining new biomarkers with
established ones is that the proposed biomarkers may seem
statistically significant in a multivariable model, i.e., have small
P values but then may not increase the prognostic ability of the
model as a whole. It can be shown statistically that in most
cases encountered in clinical prediction, very high odds ratios
or hazard ratios are required to have an effect on measures of
predictive ability (21). For a binary outcome (e.g., diseased/not
diseased), Pepe et al. (22) show that a biomarker with strong
predictive ability (21) will likely be worse even if the new assay
provides more accurate measurement. Even when changes in
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one lab compared with another.

What is Validation?

In biomarker research, validation means different things to
different people. In this issue, Chau et al. (23) discuss analytic
validation of a biomarker assay itself and describe a “fit-for-
purpose” approach in which all aspects of validation are incor-
porated. This article concerns a different form of validation:
validation of statistical models based on biomarkers. Altman
and Royston (24) distinguish between two types of validated
models: statistically validated and clinically validated. Although
both types involve evaluation of model performance, statistical
validation focuses on aspects such as goodness-of-fit, whereas
clinical validation places the prediction in context to evaluate
performance (i.e., is the prediction accurate enough for its
purpose) and might also involve considerations of costs (25).
In this paper, we focus on statistical validation.

Furthermore, we draw distinction between validating popu-
lation versus patient-level predictions. For example, the Gail
model did very well overall in predicting how many breast
cancers would occur (observed/expected ratio, 1.03), and in
general prognostic classification schemes, for example, using
stage, do reasonably well at this. However, individual pre-
dictions are more difficult and variable, consequently influen-
cing utility (26). Our focus will be on validating models at
individual level predictions.

Validation requires a comparison of model prediction with a
true outcome, which must be ascertainable or observable in the
future. For example, the true outcome (“gold standard”) might
be whether the patient has cancer, requiring an invasive
procedure to ascertain, or that a cancer will eventually recur,
requiring often lengthy follow-up to observe. In the PCPT, the
true outcome was prostate cancer status obtained via biopsy. In
the lung cancer gene expression study, the true outcome was
survival time. Some outcomes may not be ascertainable and are
thus not suited for validation. For example, Sorlie et al. (27)
gene expression data to cluster breast cancer patients into
subtypes; the true group membership of subjects in a subtype
is not ascertainable and so this outcome cannot be validated.
Validation also requires that the input data, prediction, and the
truth all be available for the subjects. Furthermore, the size of
the validation data set will be crucial to the ability to validate a
model, with larger data sets preferable.

Internal validation. In general, there are two forms of
validation. The first, internal validation, is done in the context
of an individual study, for example by splitting the study data
set into one data set to train or build the model (training set)
and one data set to test performance (test set, also called the
validation set). The appealing feature of internal validation is
its convenience, as it does not require collection of data beyond
the original study. The considerable disadvantage of the
training-testing split is reduced efficiency resulting from using
only a fraction of the data in model building. Results may
depend on the particular split of the data; thus, more
sophisticated methods than simply splitting the data into
training and testing data sets may be preferred.

For internal validation to be a valid procedure, the testing
data set must be completely untouched and no aspect of testing
data may play a role in model development. Even seemingly
innocuous uses of testing data in model building can invalidate
the internal validation attempts. For example, Potti et al. (28)
used separate training and testing gene expression data sets for
predicting response to chemotherapy. However, the authors
used the combined data to create gene clusters and used these
clusters to aid in model building using training data. Although
outcomes from the testing data were not used in model
building, their strategy allowed information to “leak” from
the testing data and invalidated the assessment of the model on
the testing data (29). An example of strict adherence to the
training/testing split is the use of an “honest broker” to provide
the model builder(s) only the required subset of data at each
step of validation. The lung cancer gene expression study used
this approach, with the broker first providing data from the
training set. After a model was chosen, he provided the gene
expression data for the testing data sets, and only after indi-
vidual predictions were made for each subject was the outcome
data for the testing data sets’ subjects made available.
An extension of the simple training-testing split is to split the
data into training and test groups a large number of times.
Commonly used examples of this approach include leave-one-
out, k-fold, and repeated random-split cross-validation. With
leave-one-out cross-validation, each observation in turn serves
as the test set with the remaining data used as the training set;
there are therefore as many testing-training splits as there are
observations. For k-fold cross-validation, the data are divided
into k subsets with each subset serving as the test set for the
remaining k-1 subsets pooled together. For repeated random-split cross-validation, the procedure of splitting the data into training and testing sets is randomly repeated many times. With cross-validation, the model is refit to each training set, evaluated on the corresponding test set, and validation results are reported as the average performance over all test sets. This allows for nearly unbiased estimates of model performance; furthermore, sample size is not sacrificed when leave-one-out cross-validation is used.

In developing the PCPT trial, 4-fold cross-validation was done. The 5,519 participants used for the analysis were split into four subsets of size 1,380, 1,380, 1,380, and 1,379, stratified by prostate cancer status so that the percentage of cancer cases ranged from 20% to 23% in each subset (1). On each possible grouping of three of the subsets, the entire model selection process used for the development of the overall PCPT Risk Model was done to yield an optimal model that was then validated on the single subset left out. The results from the four testing-training data splits were averaged to obtain a single measure of model performance.

In the lung cancer gene expression study, data came from four institutions; data from two institutions were pooled to create the training data set (N = 256) and data from the other two institutions served as separate external testing data sets (N = 104 and N = 82). There were qualitative differences in some aspects of the gene expression data for one of the external data sets, so this was viewed as a more challenging, but realistic, way to assess the performance of the model. The model was built with the training data, and an internal validation (repeated random-split cross-validation) was used to evaluate its performance. In the random splits, 200 of the 256 samples were used as the training data and the procedure was repeated 100 times. Finally, the two remaining data sets were used for an external validation (see section “External Validation”).

An advantage of cross-validation to assess performance is that the final model can be built using all the data and is more efficient than a single testing-training split. A disadvantage is that it is cumbersome and may be impossible to implement. To properly undertake cross-validation, every aspect of the model building process must be repeated independently, including all data preprocessing, selection of important variables, and estimation, as was done for the PCPT calculator and for the lung cancer gene expression study. Because aspects of model building may be subjective, automating this process can be difficult. Cross-validation permits an honest assessment of the variability of the internal validation procedure, but the method does produce a large number of different risk prediction models because the multivariable models are bound to differ among training sets, and each of these models may differ from the final model. Alternatives to cross-validation include bootstrapping methods (13, 30).

A pitfall for internal validation is that due to small sample sizes, it may be tempting to forego the split and simply evaluate the risk prediction tool with the data used to develop it. This validation will be highly biased in an overoptimistic direction, especially when a large number of biomarkers are involved. Even when an internal validation is properly done, the operating characteristics of the risk prediction tool may be overoptimistic relative to validation on a completely external data set.

**External validation.** External validation on a different data set provided by a different study circumvents these issues. Validation on heterogeneous external data sets allows for evaluation of the generalizability of the risk prediction tool to wider populations than originally reported. For example, the Gail model for breast cancer recurrence was developed using data from the Breast Cancer Detection Demonstration Project and was subsequently externally validated using data from the Breast Cancer Prevention Trial placebo arm (31). Even when external validation uses a data set that seems to arise from a familiar population, it is bound to differ due to, for example, different geographic locations or different clinical practices. It is likely that the operating characteristics of the model will be diminished compared with internal validity assessments but still may be good enough to declare the model useful. For example, external validation for the PCPT Risk Calculator yielded estimates of model performance that were lower than those initially reported using internal validation (32); more detail is given in the section “Sensitivity, Specificity, Receiver Operating Characteristic Curves, and Area Underneath the Curve.”

One caveat with external validation is that investigators only get one shot at evaluating a model on an external data set. Although it may be very instructive to use external validation results to improve the model, it is not legitimate to subsequently reevaluate the model on this external data set.

**Metrics of validation.** At the heart of the issue of validation is the statistical challenge of comparing predictions from the model with true outcomes. Although validity of a model could be couched as a yes/no determination (i.e., valid/not valid), this is not the best approach as most models would be deemed invalid, even potentially useful ones. It is useful to think in terms of degrees of validity, in particular, to assess whether a model is useful, if so, how useful, and lastly, is it as good as advertised. Whether a model performs well enough for its intended application is a question of clinical validity and is not addressed here. In terms of statistical validity, the agreement between prediction and true outcomes is usually summarized by a few measures, discussed in further sections.

**Sensitivity, specificity, receiver operating characteristic curves, and area underneath the curve.** With binary outcomes and predictions, results can be displayed in a 2 × 2 table that cross-classifies true status (e.g., subject has the disease, yes/no) with predicted status (e.g., subject is predicted to have disease, yes/no), summarizing the performance of the prognostic model. From this, one can calculate several simple measures that regularly appear in the medical literature and are thus familiar and easy to understand. Sensitivity is the proportion of the true positive outcomes (e.g., truly diseased subjects) that are predicted to be positive. Specificity is the proportion of the true negative outcomes (e.g., truly disease-free subjects) that are predicted to be negative. Because both of these measures are simple proportions, SEs and confidence intervals can easily be calculated and should be reported.

If prediction is instead on a continuous scale, such as a predicted probability or a risk score, then for any given cut-point
of the continuous scale, one could create the $2 \times 2$ table and calculate sensitivity and specificity. As part of the validation of the PCPT calculator, it was applied to a cohort from San Antonio (32). Table 2 shows how the sensitivities and specificities change as cutoffs for both PCPT risk and PSA are varied. As expected, there is a continuum of change, with the sensitivity decreasing as the specificity increases with different cutoffs.

The overall performance of the model can be summarized with a receiver operating characteristic (ROC) curve. For each possible cut-point, the resulting sensitivity and specificity are indicated as a point on a graph. This is illustrated in Fig. 1 for both PCPT risk and PSA. The overall strength of association is summarized by the area underneath the curve (AUC, often called the concordance C statistic). An AUC of 0.5 (50%) indicates no association between prediction and true outcome, and a value of 1.0 (100%) indicates perfect association. In general, AUCs below 0.6 are not considered medically useful, whereas values of 0.75 are. However, there are no absolute rules about how large the AUC must be for the predictive model to be useful, and what represents large enough will be context dependent. Finally, SEs should be calculated to capture uncertainty in the estimate of AUC; one can and should give confidence intervals. For the PCPT risk calculator in Fig. 1, the AUC is 65.5%, with 95% confidence interval 60.2% to 70.8%.

The closeness of the AUC calculated from the external data set to the AUC estimated from the data set that was used to build the model is a measure of the validity of the model. For the PCPT, the original internally validated estimate of AUC was 70.2%. The PCPT was later externally validated on the significantly younger and more ethnically diverse San Antonio population (AUC, 65.5%; $N = 446$) and on 2 observational cohorts, 1 by Johns Hopkins University (AUC, 66%; $N = 4,672$) and another at the University of Chicago (AUC, 67%; $N = 1,108$; refs. 32–34). External validation thus resulted in a reduction of $\sim 4$ percentage points from the internal validation.

ROC curves are invariant to monotonic transformations, for example, if all predictions were multiplied by a constant factor the ROC curve would remain unchanged. This property can have some unexpected results. If the continuous prediction has an interpretable scale, such as the probability of recurrence, this scale is ignored by the ROC curve; thus, it is feasible to see a high AUC when the predicted probabilities are systematically biased. For example, if risk predictions for the PCPT were all multiplied by 0.5, which would clearly give underestimates of risk, the AUC would remain unchanged compared with the original. ROC curves help describe classification performance, are useful for comparing risk tools, and can even be used to determine which biomarkers to include in the tool; however, they are only part of the assessment. A valid model should also be calibrated, that is, for example, a model of cancer risk

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<tr>
<th>PSA cutoff (ng/mL)</th>
<th>PCPT risk cutoff (%)</th>
<th>PSA and PCPT risk specificity (%)</th>
<th>Sensitivity (%), PSA</th>
<th>Sensitivity (%), PCPT risk</th>
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<tbody>
<tr>
<td>1.0</td>
<td>20.5</td>
<td>27.5</td>
<td>90.5</td>
<td>89.9</td>
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<tr>
<td>1.5</td>
<td>25.5</td>
<td>40.3</td>
<td>84.5</td>
<td>82.4</td>
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<td>2.0</td>
<td>27.5</td>
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<td>51.7</td>
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<td>36.5</td>
<td>73.8</td>
<td>39.9</td>
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<td>46.0</td>
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<td>10.0</td>
<td>59.0</td>
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should return computed risks that agree with actual observed proportions of cancer in a population of individuals with these computed risks. For example, if a model predicts that 20% of subjects will have an event within 3 years, but 40% actually do, then the model is not calibrated, regardless of how good measures such as the AUIC may be. For the PCPT calculator applied to the San Antonio cohort, the observed prostate cancer rates increased with increasing PCPT risk: 15.7%, 39.0%, 48.8%, and 100% for PCPT risk calculator values of <25%, 25% to 50%, 50% to 75%, and >75%, respectively. This shows that the PCPT risk calculator is reasonably well-calibrated.

When the true outcome is a censored event, i.e., has not yet occurred at the time of analysis, the calculation and interpretation of an ROC curve is harder. It is not legitimate to ignore the follow-up time and simply use dead or alive at last contact as a binary outcome. For the lung cancer gene expression study, the authors show how the ROC curve can be calculated for the outcome dead or alive at 3 years (9). Time-dependent ROC curves have been recently developed for situations where the true outcome in the validation data set is a censored event, such as a survival time (35).

Positive and negative predicted values. From the perspective of the patient, he or she would like to know the probability that his particular prediction is correct. For a 2 × 2 table, the positive predicted value (PPV) measures the proportion of times the true outcome is positive among those for whom it was predicted to be positive. Similarly, negative predictive value measures the proportion of true negatives out of all who tested negative. When reporting PPV and negative predictive value, one must define the population to which it applies. For example, the PPV for a predictive model in lung cancer screening is likely to be larger in a population of heavy smokers compared with a population of nonsmokers. In the context of external validation, if a predictive model claims to have a certain PPV, then one aspect of validation is to determine whether this PPV is seen in the external data set. Although PPV and negative predictive value are defined for binary outcome and prediction, they can be extended to the situation where the prediction is a continuous scale (36, 37).

When prediction is binary but the true outcome is a censored event time, it is standard practice to draw a Kaplan-Meier estimate of the time-to-event distribution for each predicted group. The PPV and negative predictive value can be read off these graphs at any desired follow-up time. A good separation between the Kaplan-Meier curves is required for the model to be useful; the relative hazard summarizes the predictive difference between groups. If the prediction has quantitative summaries associated with it, e.g., the probability of recurrence in each group, then accuracy of these claims can be assessed on external data by comparison to estimates of recurrence probability calculated from Kaplan-Meier plots (38).

When true outcome is a censored event and the prediction is a continuous score, we recommend dividing the data into at least three equal sized groups for the purpose of drawing Kaplan-Meier estimates. The reason more than two groups is preferred is because this allows an assessment of a “dose-response” relationship, which is a desirable characteristic of a predictive model. Equally sized groups are needed for objectivity. Although cut-points may not be round numbers, this avoids the pitfall of seeking out the best cut-points, which is well-known to give inflated measures of the performance (39).

Other measures. When the true outcome is binary and the prediction a probability, other measures of predictive accuracy have been suggested. The mean squared error, called the Brier score in this context (40, 41), measures the distances between true outcomes (0, no event; 1, event) and predictions (probabilities ranging from 0 to 1) for a particular test data set. A number of adaptations and generalizations of the Brier score have been proposed, including handling the situation of censored event time data (41–43). These scores are useful for comparing two competing models.

Cut-points. A common but much overemphasized practice in predictive modeling is to seek optimal cut-points or thresholds in the prediction. Although cut-points are needed in the final stages of a model to provide guidelines for medical decision making, models that provide a continuous score provide potentially useful information, particularly for subjects near the threshold. It seems unrealistic to think risk levels are truly discrete; more plausible is a continuum of risk, and for this reason, we prefer models providing a continuous score. Similarly, when subject characteristics are measured on a continuous scale, such as is often the case for biomarkers, it is implausible that there exists a fixed threshold at which risk abruptly changes. It is most efficient to keep biomarkers in the model as continuous variables if the assay measures them as such. Only at the final stage when the model is built from all the biomarkers might, it be necessary to introduce cut-points to aid in classifying people into distinct groups.

Conclusion

The challenges of validating a biomarker-based predictive model are considerable, and attention to the planned validation should be delivered as early as possible in the design phase of building the risk prediction tool. External validation using data from a completely different study provides the highest irrefutable evidence that a tool validates. The more external validations, the better, particularly when they come from more heterogeneous populations that put a stress on the generalizability of the risk tool. Internal validation is more convenient and perhaps the only option for introducing the risk tool in a timely fashion but is no substitute for external validation. For this reason, the details of the prediction tools should be published and made available for others to attempt external validation. A variety of statistical summaries, such as the AUIC, PPV, and Brier score, can be used to summarize the performance characteristics of a risk prediction tool; in practice, no one measure is enough, and the use of multiple summaries characterizing different components of prediction is recommended.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.
References


