Gene expression

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survcomp: an R/Bioconductor package for performance assessment and comparison of survival models

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ABSTRACT

Summary: The *survcomp* package provides functions to assess and statistically compare the performance of survival/risk prediction models. It implements state-of-the-art statistics to (i) measure the performance of risk prediction models; (ii) combine these statistical estimates from multiple datasets using a meta-analytical framework; and (iii) statistically compare the performance of competitive models. **Availability:** The R/Bioconductor package *survcomp* is provided open source under the Artistic-2.0 License with a user manual containing installation, operating instructions and use case scenarios on real datasets. *survcomp* requires R version 2.13.0 or higher. http://bioconductor.org/packages/release/bioc/html/survcomp.html

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1 INTRODUCTION

Building risk prediction or survival models is an important area of research, especially in cancer where gene expression signatures are used to predict risk of metastasis, response to therapy and overall survival. However, assessing the relative performance of such models is complex due to the lack of standards regarding the best criterion to use in survival analysis (Table 1; Haibe-Kains et al., 2008). Although a number of model performance estimators have been described (Cox, 1972; Graf et al., 1999; Harrel et al., 1996; Heagerty et al., 2000; Royston and Sauerbrei, 2004; Verweij and Houwelingen, 1993), they are not widely used. This is partly because these are implemented in many different R packages that use heterogeneous interfaces, which makes it difficult for the non-specialist to easily use or compare the performance of these models. Another challenge in assessing performance of expression-based prediction models is a lack of power due to small sample size. Meta-analytical methods could leverage power from the numerous microarray datasets that are now publicly available by summarizing model performance estimated in multiple-independent studies. Moreover, because they enable efficient joint analysis of multiple datasets, such an analytical framework reduces the risk of artifactual discoveries that are due to bias or confounding factors that may be present in one dataset. This is particularly important when comparing competitive risk prediction models; often authors

Name	Function in survcomp	References	
Concordance index	concordance.index	Harrel et al. (1996)	
D index	D.index	Royston and Sauerbrei (2004)	
Hazard ratio	hazard.ratio	Cox (1972)	
Brier score	sbrier.score2proba	Graf et al. (1999)	
Cross-validated	cvpl	Verweij and	
partial likelihood		Houwelingen (1993)	
Time-dependent ROC curve	tdrocc	Heagerty et al. (2000	
Kaplan–Meier curve	km.cox.plot	Kaplan and Meier (1958)	
Forestplot	forestplot.surv	Lewis and Clarke (2001)	

Table 1. Functions in *survcomp* to measure the performance of risk prediction models

claim better performance of a new model without properly assessing whether a model *significantly* outperforms its competitors. To the best of our knowledge, there is no commercial or open-source tool to enable statistical comparison of risk prediction models in a meta-analytical framework.

To address these issues, we developed a new Bioconductor package, *survcomp*, which implements several performance criteria for risk prediction models (Table 1), together with metaanalytical methods that enable combination of performance estimations from multiple-independent datasets [fixed- and random-effects models (Cochrane, 1954); ?combine.est], and statistical comparison of predictions between competitive models (?cindex.comp for the concordance index). The concordance index as described by Harrel *et al.* (1996) and implemented in *survcomp* may be sensitive to the study-specific censoring distribution, therefore we are working to implement a modified concordance index by Uno *et al.* (2011) that avoids this problem and which should be available in the next release of *survcomp*.

Although the performance criteria that are implemented in *survcomp* are mostly available in other R packages (except the D index which, to the best of our knowledge, is only in *survcomp*), our package provides a common interface to facilitate easy use of all these estimators. Moreover, with the exception of *ipdmeta* by Broeze *et al.* (2009) and *survJamda* by Yasrebi (2011), few R packages perform meta-analysis of survival data. *survcomp* provides a uniform interface to simplify the use of performance assessment and statistical comparison of risk prediction models, and provides new R functions to statistically compare these in

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Fig. 1. Forestplots representing the prognostic value of (a) NPI, (b) AURKA, (c) GGI and (d) GENIUS estimated by the concordance index in five independent breast cancer datasets [none of these datasets were used to train the classifiers (a)–(c), duplicated patients were removed what results in a combined dataset of 722 patients]. The blue square and horizontal line represent the concordance index and its 95% confidence interval which is clipped at 0.4 and 0.9 (represented by an arrow). The black rhombus is the overall meta-estimate from the combined five datasets. The greater the concordance index, the more prognostic the risk prediction model. The vertical red bar represents the concordance index of random risk predictions.

a meta-analytical framework. It is worth noting that the aim of *survcomp* is to provide efficient computation of several performance estimates (Table 1) and not to implement a full validation framework. Proper validation using a fully independent test dataset is of utmost importance to avoid overoptimistic results (Jelizarow *et al.*, 2010) and frameworks for cross-validation, multiple random splits or a single split into training/validation datasets are implemented in several existing R packages including *peperr* (Porzelius *et al.*, 2009).

We illustrate the functionality of *survcomp* in a case study that investigates prognosis in breast cancer. In most studies, this is a difficult task due to the (generally) small sample size. Yet despite a large number of published studies, data heterogeneity, both in terms of data sources and microarray technologies, have limited the effectiveness of their joint analysis. Here we apply *survcomp* functions to statistically compare the prognostic value of a widely used clinical model, a single proliferation gene and two published gene signatures. The results suggest that the multi-gene signatures are not always superior to standard clinical models or to a simple single gene model.

2 CASE STUDY

The prognostic ability of gene expression of AURKA, a single proliferation-related gene, was compared with the Nottingham Prognostic Index (NPI, Galea *et al.*, 1992) clinical model for prognosis, and to risk prediction scores from two published multigene prognostic signatures; GGI (Sotiriou *et al.*, 2006) and GENIUS (Haibe-Kains *et al.*, 2010). The NPI, GGI and GENIUS risk scores were calculated using the Bioconductor package *genefu*. Each score was computed in five publicly available datasets described in the Supplementary Material.

To compare the prognostic ability of these four different risk prediction models, we estimated the concordance index for each model in each dataset separately and used the function combine.est to compute the corresponding overall meta-estimate using the well-established random-effects model approach (Cochrane, 1954). As can be seen in Figure 1, although the performance varies between datasets, all models yielded highly significant overall prognostic value (high-risk predictions represent patient with poor survival, concordance index > 0.5, one-sided P <

 Table 2. Statistical performance comparison for the risk prediction models used in our case study

Models	NPI	AURKA	GGI	GENIUS
NPI		0.25	0.37	0.91
AURKA	0.75		0.70	0.98
GGI	0.63	0.30		0.97
GENIUS	0.09	0.02	0.033	

P-values are computed using a one-sided paired Student's *t*-test to test whether risk prediction models in rows are better than the ones in columns.

0.001, see Supplementary Material for R code). AURKA, the single proliferation gene, was the worst predictor of survival (concordance index of 0.64), whereas GENIUS, the risk prediction model taking into account the breast cancer molecular subtypes, was the best (concordance index of 0.69). The continuous risk prediction of NPI, the traditional clinical model which combines nodal status, tumor size and histological grade, yielded a relatively high concordance index (concordance index of 0.66).

To identify the best risk prediction model(s), we statistically compared their concordance indices using the function cindex.comp.meta (Table 2). Concurring with Haibe-Kains *et al.* (2010) we observed that GENIUS outperforms AURKA, GGI and NPI (uncorrected P < 0.10; Table 2). However, when *P*-values are corrected for multiple testing (Holm's method), no *P*-value remains significant suggesting that a larger meta-analysis is required to definitively claim the superiority of one classifier over another. This also suggests that prognostic clinical models such as NPI are still extremely competitive compared with more complex gene signatures. Repeating this analysis using performance criteria in *survcomp* other than the concordance index, including the D index or hazard ratio (Table 1), leads to similar conclusions (examples are provided in the package user manual and documentation).

3 CONCLUSION

The R/Bioconductor package *survcomp* provides a uniform interface to an extensive set of performance assessment and statistical comparison methods for survival/risk prediction. It allows scientists to easily implement large comparative studies integrating multiple independent datasets while providing statistical tools to identify the best model(s) as supported by the data under study.

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