Effect of machine learning methods on predicting NSCLC overall survival time based on Radiomics analysis

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Abstract

Background: To investigate the effect of machine learning methods on predicting the Overall Survival (OS) for non-small cell lung cancer based on radiomics features analysis.

Methods: A total of 339 radiomic features were extracted from the segmented tumor volumes of pretreatment computed tomography (CT) images. These radiomic features quantify the tumor phenotypic characteristics on the medical images using tumor shape and size, the intensity statistics and the textures. The performance of 5 feature selection methods and 8 machine learning methods were investigated for OS prediction. The predicted performance was evaluated with concordance index between predicted and true OS for the non-small cell lung cancer patients. The survival curves were evaluated by the Kaplan-Meier algorithm and compared by the log-rank tests.

Results: The gradient boosting linear models based on Cox’s partial likelihood method using the concordance index feature selection method obtained the best performance (Concordance Index: 0.68, 95% Confidence Interval: 0.62~0.74).

Conclusions: The preliminary results demonstrated that certain machine learning and radiomics analysis method could predict OS of non-small cell lung cancer accuracy.

Keywords: Overall survival, Non-small cell lung cancer, Machine learning, Radiomics analysis

Background

Lung cancer is the leading cause of cancer-related deaths worldwide [1]. Lung cancer could be clinically divided into several groups: 1) the non-small cell lung cancer (NSCLC, 83.4%), 2) the small cell lung cancer (SCLC, 13.3%), 3) not otherwise specified lung cancer (NOS, 3.1%), 4) Sarcoma lung cancer (0.2%), and 5) other specified lung cancer (0.1%) [2]. The ability to predict clinical outcomes accurately is crucial for it allows clinicians to judge the most appropriate therapies for patients.

Radiomics analysis can extract a large number of imaging features quantitatively, which could offer a cost-effective and non-invasive approach for individual medicine [3–5]. Several studies have shown the predictive and diagnostic ability of radiomics features in different kinds of cancers using various medical imaging modalities, such as PET [6–8], MRI [9] and CT [4, 10, 11]. It is also demonstrated that the radiomic features are associated with the overall survival. Besides, these associations can be used to establish positive predictive models.

Machine-learning (ML) can be resumptively defined as the computational methods utilizing data/experience to obtain precise predictions [12]. The ML method can first learn laws from the data and then establish accuracy and efficiency prediction model based on these laws automatically. Moreover, an appropriate model is essential for the success use of radiomics. Hence, it is crucial to compare the performance of different ML models for clinical biomarkers based on radiomics analysis. Besides, appropriate feature selection methods should be applied first for the high-throughput radiomics features who may cause serious overfitting problems.

In this study, we investigated the effect of 8 ML and 5 feature selection methods on predicting OS for non-small cell lung cancer based on radiomics analysis. The effectiveness of ML and feature selection methods
on the prediction of OS were evaluated utilizing the concordance index (CI) [6, 13–16].

Methods
Data acquisition
The data used in this study was obtained from the ‘NSCLC-Radiomics’ collection [4, 17, 18] in the Cancer Imaging Archive which was an open access resource [19]. All the NSCLC patients in this data set were treated at MAASTRO Clinic, the Netherlands. For each patient, manual region of interest (ROI), CT scans and survival time (including survival status) were available. All the ROIs in this data set were the 3D volume of the gross tumor volume (GTV) delineated by a radiation oncologist.

Prediction process
The flow chart of the prediction process [20, 21] for all the ML methods in this study was outlined in Fig. 1. The performance of each ML and feature selection methods for the 283 NSCLC patients were evaluated using the cross-validation (CV) method (3-CV in this study). For each CV process, the total patients were divided into three folds, in which two folds (training fold) for training the machine learning model and the third (validation fold) for evaluating the model.

For each training fold, the training algorithm required both the training inputs (for prediction) and the prediction targets (for validation) data. The training inputs referred to the selected radiomics features, while the prediction targets referred to the OS of the patients. The radiomics features were first extracted from the images and then selected (dimension reduction) using the filter based feature selection methods to reduce the risk of overfitting. Finally, the selected features would be used to optimize and train all the ML models. In this study, the Bayesian optimization method was applied to determine the optimal parameters [22].

For each validation fold, the corresponding selected radiomics features were first extracted from the images and then transferred into the trained model. Finally, the prediction OS would be used to evaluate the goodness of each model.

Image pre-processing and Radiomics features extraction
Prior to extracting the radiomics features, we fixed the bin number (32 bins) of all the pre-treatment CT scans to discretize the image intensities. It should be noted that the original voxels for the images were used in this study. Then, the radiomics features were automatically extracted from the GTV region of the CT images by our in-house developed radiomics image analysis software and the Wavelet toolbox based on the Matlab R2017a (The Mathworks, Natick, MA). Total 43 unique quantitative features in 4 categories (Fig. 2) were extracted:

1) Intensity features: to describe the shape characteristics of the CT volume’s gray-level intensity histogram, i.e., a probability density function (PDF) of gray-level distribution.

2) Fine texture features: to describe the high-resolution heterogeneity in the ROI. These features were derived from the ROI’s Gray-Level Co-Occurrence Matrix (GLCOM), a joint PDF that measures the frequency of co-occurring adjacent voxel pairs having the same gray-scale intensity at a given direction [23].

Fig. 1 The flow chart of predicted process for each ML method. (I) Dividing total data into three folds using the cross validation method. (II) Training each ML model using the selected radiomics features of the training fold. (III) Validating the prediction performance of each ML model on the validation fold.
3) Coarse texture features: to describe the low-resolution heterogeneity in the ROI. These features were calculated from the ROI’s Gray-Level Run Length Matrix (GLRLM), a joint PDF that measures the size of a set of consecutive voxels with the same grayscale intensity at a given direction [24].

4) Morphological features: to describe the morphological characteristics of the ROI [25].

Here, the first category and the following two (second and third) categories required the intensity histogram and textural image processing steps, respectively. Both the above two image processing steps and the 43 radiomics features used in this study matched benchmarks of the Image Biomarker Standardization Initiative (IBSI) [26].

Moreover, these radiomics features were also extracted from different wavelet decompositions of the original CT image by a three levels wavelet transformation [27, 28]. However, the morphological features weren’t extracted from the images with the wavelet decompositions for the wavelet transformation didn’t have effect on these features. Hence in total, 339 features were extracted for each patient in this study.

Features selection and machine learning methods
Pearson’s (PCC) [29], Kendall’s (KCC), Spearman’s linear correlation coefficient (SCC) [31], Mutual information (MI) [32] and CI [15] were used as the filter based feature selection methods to reduce the dimensions of radiomics features in this study. In order to make sure the reliability of the selected features, we repeated each feature selection process 100 times using the bootstrap samples of each training fold and recorded the selected feature subset each time. Then, we selected the most frequently selected radiomics features as the final features which were used to train the ML models [6]. In this study, the first four feature selected methods (PCC, KCC, SCC and MI) were implemented using the Matlab R2017a and the following one method (CI) was implemented using the R software 3.5.1. All the feature selection methods would be performed on each training fold.

The effect of 8 ML methods were investigated in this study, including Cox proportional hazards model (Cox) [33], gradient boosting linear models based on Cox’s partial likelihood (GB-Cox) [34], gradient boosting linear models based on CI’s partial likelihood (GB-Cindex) [34], Cox model by likelihood based boosting (CoxBooxt) [35], ...
bagging survival tree (BST) [36], random forests for survival model (RFS) [37], survival regression model (SR) [38] and support vector regression for censored data model (SVCR) [39, 40]. All the machine learning methods were implemented on each training fold using the R software 3.5.1. The specifics of the packages for each feature selection and ML method were showed in the Table 1. Besides, the descriptions of each feature selection and ML method could be found in the Additional file 1: Supplementary A and B, respectively.

Parameters tuning
For each ML method, the parameters were selected from the combination of parameters that produced the best performance using the three-fold CV on each training fold. Similar procedures were implemented in Brungard et al. [41] and Heung B et al [42].

The range of parameters used in this study was showed in Table 1. The GB-Cox, GB-Cindex, SVCR and SR methods just required one parameter to tune while the Cox method did not require parameterization. The complex models, such as the BTS and RFS, were time consuming for tuning parameters. The parameters from all of these models, such as the average terminal node size of forest and the number of trees for the RFS model, the minimum number of observations that must exist in a node (Minsplit) and the number of trees for BST, made up a large range of parameter permutation and combination choices. It should be noted that the feature number selected by the feature selection methods were also used as a tuning parameter (range [3, 29]) for all the ML methods.

Evaluation methods
CI with confidence interval (CFI) based on bootstrapping technique (the number of bootstrap samples was 2000 in this study) was used to assess the performance of difference ML methods on the merged validation fold (merged all the three validation folds). The percentage of CFI was 95% in this study. A nonparametric analytical approach method proposed by Kang L et al. [43] and the z-score test method were used to compare the significance between pairs of machine learning algorithms for each validation fold. Besides, the survival curves were evaluated by the Kaplan-Meier algorithm and compared by the log-rank tests [44] for each validation fold.

Table 1 The specifics of the packages for each feature selection and machine learning method

| Methods | Software       | Packages     | Website Links               |
|---------|----------------|--------------|-----------------------------|
| PCC     | SML toolbox    | corr         | https://ww2.mathworks.cn/help/stats/corr.html |
| KCC     |                |              |                             |
| SCC     |                |              |                             |
| MI      | MTToolbox      | mi           | https://github.com/Craigacp/MIToolbox |
| CI      | Hisc           | rcorr.cens   | https://github.com/harrelfe/Hmisc |
| Cox     | survival       | coxph        | https://github.com/therneau/survival |
| GB-Cox  | mboost         | mboost       | https://github.com/boost-R/mboost |
| GB-Cindex| mboost        | mboost       | https://github.com/boost-R/mboost |
| CoxBoost| CoxBoost       | CoxBoost     | https://github.com/binderh/CoxBoost |
| BST     | ipred          | bagging      | https://github.com/cran/ipred |
| RFS     | randomForestSRC| rfsrc       | https://github.com/kogalur/randomForestSRC |
| SR      | survival       | survreg      | https://github.com/therneau/survival |
| SVCR    | survivalsvm    | survivalsvm  | https://git-hub.com/imbs-hl/survivalsvm |

Patients on each validation fold were divided into two groups (low- and high- risk group) based on the

Results
Figure 3 depicted the performance of ML (in rows) and feature selection methods (in columns) on the merged validation fold. Besides, the maximum CI with confidence interval for each ML method on the merged validation fold was showed in Table 2. The GB-Cox method using the CI feature selection method obtained the best performance (CI: 0.682, 95% CFI: [0.620, 0.744]). However, the CoxBoost method using CI feature selection method also obtained a favorable performance (CI: 0.674, 95% CFI: [0.615, 0.731]). We found only the above mentioned two prediction method’s CIs were close. Hence, we just calculated the p-value using the z-test between the above two methods. The p-value of CI between these two methods was 0.5, indicating that the difference of prediction performance between these two methods wasn’t significant. The values selected for the hyper-parameters mentioned in Table 3, as well as the number of selected features on each validation fold could be found in the Additional file 1: Supplementary C.
predicted risk of each radiomics model at the cut-off value. The cut-off value utilized for stratification was the median of each training fold which would be applied to the corresponding validation fold unchanged. Then, the Kaplan-Meier and log-rank tests methods were used to evaluate and compare the survival curves for each validation fold, respectively. Among all the ML methods, the GB-Cox method with the CI feature selection method obtained the best stratified result on the 3 CV folds (Fig. 4). Besides, the p-value of the CoxBoost method with the PCC feature selection method was also significant for each validation fold. The heatmap of p-values on each validation fold for all the ML methods was showed in the Additional file 1: Supplementary D.

Discussion
Several previous studies have compared the prediction performance of the ML models based on the radiomics analysis. Parmar C et al. [11] identified that three classifiers, included Bayesian, random forest (RF) and nearest neighbor, showed high OS prediction performance for the head and neck squamous cell carcinoma (HNSCC). Parmar C et al. [17] also evaluated the effect of ML models (classifiers) on the OS prediction for NSCLC patients and found that the random forest method with maximally selected rank statistics and the model based on boosting trees using CI methods with Spearman feature selection method got the best prediction performance for the loco-regional tumor control. Besides, the survival regression model based on the Weibull distribution, the GB-Cox and the GB-Cindex methods with the random feature selection method achieved the highest prediction performance for the OS. In this study, the effect of 8 ML models and five feature selection methods based on radiomics feature analysis were investigated to predict the time-to-event data (OS) of non-small cell lung cancer. In general, the GB-Cox method obtained the best predictive performance in the systematic evaluation on the merged validation

![Fig. 3](image_url)

The performance of feature selection and machine learning methods on the merged validation fold

| Methods | FS | Maximum CI | CFI of Maximum CI |
|---------|----|------------|-------------------|
| GB-Cox  | CI | 0.682      | [0.620, 0.744]    |
| CoxBoost| CI | 0.674      | [0.615, 0.731]    |
| Cox     | MI | 0.646      | [0.578, 0.714]    |
| GB-Cindex| SCC | 0.357      | [0.290, 0.423]    |
| RFS     | PCC | 0.627      | [0.558, 0.695]    |
| SR      | MI | 0.380      | [0.310, 0.452]    |
| BST     | SCC | 0.385      | [0.318, 0.450]    |
| SVCR    | KCC | 0.405      | [0.341, 0.470]    |

FS feature selection method

Table 3 The range of parameter tuning

| Methods    | Parameters                        | Range of Parameters |
|------------|-----------------------------------|---------------------|
| GB-Cox     | Number of boosting steps          | [1, 500]            |
| GB-Cindex  | Number of boosting steps          | [1, 500]            |
| Coxboost   | Number of boosting steps          | [1, 500]            |
| BST        | Minsplit                          | [1, 10]             |
|            | Number of trees                   | [1, 500]            |
| RFS        | Average terminal node size of forest | [1, 10]         |
| SR         | Assumed distribution              | Weibull, Gaussian, Exponential |
| SVCR       | Parameter of regularization       | [0.01, 1]           |
fold. However, the CoxBoost methods with certain feature selection method also showed comparable positive performance compared with the GB-Cox method. Hence, we thought a wide range of ML methods have the potential to be effective radiomics analysis tools. Besides, a significant difference for OS prediction on each validation fold was found between the low- and high- risk groups using the GB-Cox and CoxBoost methods, which showed the clinical potential of ML methods on the OS prediction.

As shown in Fig. 3, almost all of the ML methods using the KCC feature selection method didn’t obtain a positive result. This indicated that the feature selection method was also important for the performance of OS prediction. Sometimes, the effect of feature selection methods was even more obvious than the ML models. A large panel of feature selection methods had been used for data mining of high-throughput problems [45, 46]. In general, the feature selection methods would be divided into three categories: the filter based, the wrapper based and the embedded methods. In this study, we only investigated five different filter based methods because this kind of methods were not only less prone to overfitting but also more efficient in computation than other two methods [45, 46]. Moreover, the filter based methods were more independent than the wrapper and embedded methods, which could increase the fairness of ML methods comparison.

Some previous studies [4, 5] have shown the potential clinical utility of the prognostic models based on radiomics analysis. This study could be a crucial supplementary reference for the use of prognostic models based on radiomics analysis because we compared a large number of machine-learning methods for the OS prediction of the NSCLC cancer. Such a comparison would be helpful in the selection of the optimal ML methods for OS prediction based on radiomics analysis.

Conclusion
The preliminary results demonstrated that certain machine learning and radiomics analysis method could predict OS of non-small cell lung cancer accuracy.

Additional file

**Additional file 1**: Supplementary A: Feature selection methods. Supplementary B: Machine learning methods. Supplementary C: The values selected for the hyper-parameters on each validation fold. Supplementary D: P-values of the log-rank test for all the feature selection and ML methods on each validation fold. (PDF 625 kb)

Abbreviations
BST: Bagging survival tree; CFI: Confidence interval; CI: Concordance index; Cox: Cox proportional hazards model; CoxBoost: Cox model by likelihood based boosting; CT: Computed tomography; CV: Cross-validation; GB-Cindex: gradient boosting linear models based on concordance index; GB-GCox: gradient boosting linear models based on Cox’s partial likelihood; GLCOM: Gray-level co-occurrence matrix; GLRLM: Gray-level run length matrix; GTV: Gross tumor volume; HNSCC: head and neck squamous cell carcinoma; KCC: Kendall’s correlation coefficient; MI: Mutual information;
ML: Machine-learning; NSCLC: Non-small cell lung cancer; OS: Overall survival; 
PCC: Pearson’s correlation coefficient; PDF: Probability density function; 
RFS: Random forests for survival model; ROI: Region of interest; 
SCC: Spearman’s linear correlation coefficient; SCLC: Small cell lung cancer; 
SR: Survival regression model; SVCR: Support vector regression for censored 
data model

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Availability of data and materials
The datasets used in this study are available.

Authors’ contributions
WS and MJ designed the methodology. WS and PC written the program. WS 
written the manuscript. DJ, MJ and FY reviewed the manuscript. All authors 
read and approved the final manuscript.

Ethics approval and consent to participate
Data collection was approved by the local IRB.

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References
1. Bhattacharjee A, Richards WG, Staunton J, Li C, Monti S, Vasa P, et al. 
Classification of human lung carcinomas by mRNA expression profiling 
reveals distinct adenocarcinoma subclasses. Proc Natl Acad Sci U S A. 2001; 
98(24):13790–5.
2. Hoadley N, Noone AM, Krapcho M, et al. SEER Cancer statistics review, 
1975–2012. Seer.cancer.gov/cs/1975_2012/ Bethesda, MD: National Cancer 
Institute; 2013.
3. Gillies RJ, Kinahan PE, Hricak H. Radiomics: images are more than pictures, 
they are data. Radiology. 2015;278(2):563–77.
4. Aerts HJ, Velazquez ER, Leijenaar RT, et al. Decoding tumour phenotype by 
noninvasive imaging using a quantitative radiomics approach. Nat Commun. 
2014;5:4006.
5. Vallieres M, Zwanenburg A, et al. Responsible radiomics research for faster 
clinical translation. J Nucl Med. 2018;59:189–93.
6. Cui Y, Song J, Pollom E, et al. Quantitative analysis of 18F-
Fluorodeoxyglucose positron emission tomography identifies novel 
prognostic imaging biomarkers in locally advanced pancreatic cancer 
patients treated with stereotactic body radiation therapy. Int J Radiat Oncol Biol 
Phys. 2016;96(1):102–9.
7. Lampropoulos M, Stathakis P, Starmans MH, et al. Predicting outcomes in 
radiation oncology–multifactorial decision support systems. Nat Rev Clin 
Oncol. 2013;10(1):27–40.
8. Chen HH, Su W, Hsieh W, Wu Y, Lin F. Summation of F18-FDG uptakes on 
PET/CT images predicts disease progression in non-small cell lung cancer. 
Int J Radiat Oncol. 2010;78(3):5504.
9. Tiwari P, Kurhanewicz J, Madabhushi A. Multi-kernel graph embedding for 
detection, Gleason grading of prostate cancer via MRI/MRS. Med Image 
Anal. 2013;17(2):219–35.
10. Ahmad C, Christian D, Matthew T, Bassam A. Predicting survival time of 
lung cancer patients using radiomic analysis. Oncotarget. 2017;8(61): 
104393–407.
11. Parmar C, Grossmann P, et al. Radiomic machine-learning classifiers for 
prognostic biomarkers of head and neck cancer. Front Oncol. 2015;5:272.
12. Mohri M, Rostamizadeh A, Talwalkar A. Foundations of machine learning. 
Ch. 1, 1–3, MIT press, 2012.
13. Leger S, Zwanenburg A, et al. A comparative study of machine learning 
methods for time-to-event survival data for radiomics risk modelling. Sci 
Rep. 2017;7:13206.
14. Harrell FE Jr, Lee KL, Mark DB. Tutorial in biostatistics: multivariable 
probabilistic models: issues in developing models, evaluating assumptions and 
adequacy, and measuring and reducing error. Stat Med. 1996;15(4):361–87.
15. Newson R. Confidence intervals for rank statistics: Somers’ D and extensions. 
Stata J. 2006;6(3):309–34.
16. Harrell FE. Regression modeling strategies: with applications to linear 
models, logistic regression, and survival analysis. New York: Springer science 
& business media; 2001.
17. Parmar C, Grossmann P, et al. Machine learning methods for quantitative 
Radiomic biomarkers. Sci Rep. 2015;5:13087.
18. Aerts HJ, Rios V, et al. Data from NSCLC-Radiomics. Cancer Imaging Archive. 2015.
19. Clark K, Vendt B, Smith K, et al. The Cancer imaging archive (TCA): 
maintaining and operating a public information repository. J Digit Imaging. 
2013;26(6):1045–57.
20. Collins GS, Reitsma JB, et al. Transparent reporting of a multivariable 
prediction model for individual prognosis or diagnosis (TRIPOD): the TRIPOD 
statement. Ann Intern Med. 2015;162:55.
21. Moons KGM, Altman DG, et al. Transparent reporting of a multivariable 
prediction model for individual prognosis or diagnosis (TRIPOD): 
explanation and elaboration. Ann Intern Med. 2015;162:W1.
22. Snoek J, Larochelle H, Adams RP. Practical Bayesian optimization of machine 
learning algorithms. Adv Neural Inf Proces Syst. 2012;2:2951–9.
23. Haralick RM, Shanmugam K. Textural features for image classification. IEEE 
Trans Syst Man Cybern. 1973;3(6):610–21.
24. Tang X. Texture information in run-length matrices. IEEE Trans Image 
Process. 1998;7(11):1602–9.
25. Guo W, et al. Prediction of clinical phenotypes in invasive breast carcinomas 
from the integration of radiomics and genomics data. J Med Imaging (Bellingham). 
2015;2(4):041007.
26. Zwanenburg A, Leger S, Valléeys M, Löck S. Image biomarker standardization 
initiative arXiv161207003. 2016.
27. Selsnick I. The double density DWT wavelets in signal and image analysis: 
from theory to practice. Norwell: Kluwer Academic Publishers; 2001.
28. Selsnick I, Baraniuk RG, Kingsbury KG. The dual-tree complex wavelet 
transform. IEEE Signal Processing Mag. 2005;22(6):123–51.
29. Karl P. Notes on regression and inheritance in the case of two parents. Proc 
R Soc London. 1895;58(1895):240–8.
30. Kendall M. A new measure of rank correlation. Biometrika. 1938;30(1–2):81–9.
31. Jerome LM, Arnold DW. Research design and statistical analysis 2nd. 
Mahwah: Lawrence Erlbaum; 2003.
32. Pocock A, Zhao MJ, Luján M. Conditional likelihood maximisation: a unifying 
framework for information theoretic feature selection gavin brown. J Mach 
Learn Res. 2012;13:27–66.
33. Andersen P, Gill R, Cox’s regression model for counting processes, a large 
sample study. Ann Stat. 1982;10:1100–20.
34. Hofner B, May R, Robinzonov N, Schmid M. Model-based boosting in R: a 
hands-on tutorial using the R package mboost. Comput Stat. 2014;29:3–35.
35. Binder H, Allignol A, Schumacher M, Beyersmann J. Boosting for 
high-dimensional time-to-event data with competing risks. Bioinformatics. 
2009;25:890–6.
36. Hothorn T, Lauzen B, Benner A, Radespiel-Troeger M. Bagging survival trees. 
Stat Med. 2004;23(1):77–91.
37. Ishwaran H, Kogalur UB, Blackstone EH, Lauer MS. Random survival forests. 
Ann Appl Stat. 2008;2:841–60.
38. Kalbflesch JD, Prentice RL. The statistical analysis of failure time data. New 
York: Wiley; 2002.
39. Van Belle V, Pelcmans K, et al. Improved performance on high-dimensional survival 
data by application of survival-SVM. Bioinformatics (Oxford). 2011;27:87–94.
40. Van Belle V, Pekmans K, et al. Support vector methods for survival analysis: a comparison between ranking and regression approaches. Artif Intell Med. 2011;53:107–18.

41. Brungard CW, Boettinger JL, et al. Machine learning for predicting soil classes in three semi-arid landscapes. Geoderma. 2015;239-240:83–83.

42. Heung B, Bulmer CE, Schmidt MG. Predictive soil parent material mapping at a regional scale: a random forest approach. Geoderma. 2014;214-215:151-154.

43. Kang L, Chen W, Petrick NA, Gallas BD. Comparing two correlated C indices with right-censored survival outcome: a one-shot nonparametric approach. Stat Med. 2014;34(4):685–703.

44. Royston P, Altman DG. External validation of a cox prognostic model: principles and methods. BMC Med Res Methodol. 2013;13:33.

45. Bolón-Canedo V, Sánchez-Maroño N, et al. Review of microarray datasets and applied feature selection methods. Inform Sciences. 2014;282(20):111–135.

46. Guyon I, Elisseeff A. An introduction to variable and feature selection. J Mach Learn Res. 2003;3(6):1157–82.