Synergistic combination of a topologically invariant imaging signature and a biomarker for the accurate prediction of symptomatic radiation pneumonitis before stereotactic ablative radiotherapy for lung cancer: A retrospective analysis

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Abstract

Objectives

We aimed to explore the synergistic combination of a topologically invariant Betti number (BN)-based signature and a biomarker for the accurate prediction of symptomatic (grade ≥ 2) radiation-induced pneumonitis (RP+) before stereotactic ablative radiotherapy (SABR) for lung cancer.

Methods

A total of 272 SABR cases with early-stage non-small cell lung cancer were chosen for this study. The occurrence of RP+ was predicted using a support vector machine (SVM) model trained with the combined features of the BN-based signature extracted from planning computed tomography (pCT) images and a pretreatment biomarker, serum Krebs von den Lungen-6 (BN+KL-6 model). In all, 242 (20 RP+ and 222 RP–(grade 1)) and 30 cases (8 RP+ and 22 RP–) were used for training and testing the model, respectively. The BN-based features were extracted from BN maps that characterize topologically invariant heterogeneous traits of potential RP+ lung regions on pCT images by applying histogram- and texture-based feature calculations to the maps. The SVM models were built to predict RP+ patients with a BN signature that was constructed based on the least absolute shrinkage and selection operator logistic regression model. The evaluation of the prediction models was performed based on the area under the receiver operating characteristic curves (AUCs) and accuracy in the test. The performance of the BN+KL-6 model was compared to the
performance based on the BN, conventional original pCT, and wavelet decomposition (WD) models.

**Results**

The test AUCs obtained for the BN+KL-6, BN, pCT, and WD models were 0.825, 0.807, 0.642, and 0.545, respectively. The accuracies of the BN+KL-6, BN, pCT, and WD models were found to be 0.724, 0.708, 0.591, and 0.534, respectively.

**Conclusion**

This study demonstrated the comprehensive performance of the BN+KL-6 model for the prediction of potential RP+ patients before SABR for lung cancer.

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**1. Introduction**

Stereotactic ablative radiotherapy (SABR) is a non-invasive treatment for early-stage non-small cell lung cancer (NSCLC). Previous studies have shown that the overall survival and progression-free survival of patients treated with SABR are comparable to those of lobectomy, although it is a much less invasive procedure [1, 2]. However, SABR can cause symptomatic (grade ≥ 2) radiation-induced pneumonitis (RP+) in 28% of patients [3, 4]. Although most cases of RP are manageable, a few cases are severe, and there is a risk of mortality [3]. This raises the demand for predicting the occurrence of RP+ before SABR to support radiation oncologists in decision-making for radiotherapy.

An antigen Krebs von den Lungen-6 (KL-6), mucin-like glycoprotein, has been recognized as a biomarker of pulmonary epithelial cell injury [5–8]. Besides it is widely known for its association with the activity of interstitial pneumonia (IP), a number of studies reported the usefulness of the KL-6 for the prediction of RP+ prior to SABR [5–7, 9]. They mentioned that patients with elevated KL-6 levels showed the tendency to have RP+ after SABR. Therefore, prescreening of KL-6 is suggested to decide the strategy for SABR in lung cancer patients [5–7]. However, since KL-6 is associated with RP+ as well as the existence of lung cancer itself [10, 11], the predictive performance of pretreatment KL-6 may not be sufficient (S1 Table).

The potential of planning computed tomography (pCT) image signatures in the prediction of RP status after SABR has been explored in previous radiomics studies [12, 13]. Hirose et al. utilized histogram and texture features extracted from pCT and wavelet decomposition (WD) images, which have been widely used in radiomics studies, to construct the image signature and prediction model using the least absolute shrinkage and selection operator logistic regression (LASSO-LR) [13]. Moran et al. attempted to develop a LR model for the classification of the RP status based on the image features extracted from the pCT as well as the post-SABR diagnostic CT images [12]. Although their model demonstrated the feasibility for RP prediction, their area under the receiver operating characteristics (ROC) curve (AUC) could not reach a value higher than 0.76 [12, 13]. The prediction performance is not sufficient for clinical settings. Further, to our knowledge, none of these studies have investigated the impact of the combination of image signature and KL-6 in RP prediction.

Pre-existing pulmonary diseases (prePDs) have been recognized as predictive factors for the occurrence of RP after radiation treatments, including SABR [14–18]. The prePDs associated with RP can be categorized into interstitial pneumonia (IP) and chronic obstructive
It has been reported that patients with the IP had a higher risk of RP+ [15, 16], whereas patients with COPD had a lower risk of RP+ [14, 17, 18]. Importantly, IP and COPD cause structural changes of lung regions, which have been observed in pathological images [19–21]. Consequently, those microstructural changes of lung regions also appear in CT images. IP increases X-ray attenuation as a result of dense fibrotic remodeling of the lung structure [19, 20, 22]. The remodeling results in ground-glass opacities that derive spatially heterogeneous pixel value distributions on CT images. In contrast, the COPD molds lower-attenuation areas, which can be observed as cavitation on the CT images, and are caused by a lack of alveolar surrounding airways [21, 23]. These prePDs could cause pulmonary structural alteration and lead to heterogeneous and/or cavitated lung textures caused by low and/or high attenuation areas on CT images.

We have developed a novel topological image processing technique utilizing Betti numbers (BNs), which represent the number of connected components (b0) and holes (b1), to preserve the intrinsic heterogeneous patterns and cavitation of objects. BN-based image features have demonstrated feasibility in the quantification of tumor heterogeneity and cavitation for prognostic prediction [24] or identification of epidermal growth factor receptor mutations among patients with NSCLC [25]. Further, it has shown robustness against variability in imaging parameters and patient populations [24, 25]. Therefore, we hypothesized that RP pulmonary structural patterns with prePDs on pCT images can be characterized using the versatile topological image processing technique based on BNs. Consequently, the BN signature could be a promising predictive factor for RP before SABR.

Thus, we aimed to develop a comprehensive image signature based on BNs to accurately predict the occurrence of RP+ using pCT images before SABR for lung cancer. Furthermore, we investigated its synergistic relationship with KL-6 to consider the practical use of the BN-based signature in combination with this biomarker. This is the first study exploring the feasibility of a combination of the robust image features based on the BN and the biomarker KL-6. The accurate prediction of the RP+ could potentially guide physicians to determine the optimal dose prescription to avoid severe RP+ in personalized SABR.

2. Materials and methods

2.1. Clinical cases

A total of 272 patients who underwent SABR for NSCLC were retrospectively enrolled in this study. Patient data for the training dataset were randomly selected during the period between August 2003 and July 2013. As for the test dataset, we collected all the available cases with the RP+, and randomly selected the grade 1 RP (RP–) cases in the period between April 2014 and March 2018, so that the number of cases in both groups was not too different from each other. By doing so, we assumed that the bias to the majority class can be avoided in the evaluation of the performance of the proposed models. [26–29]. When the number of cases in either major or minor class is too small (or high), the model would have a high pseudo accuracy by predicting all the cases as belonging to the major class; but when it comes to the minor class, the performance deteriorates [26–29]. Therefore, the number of RP–was reduced to avoid inappropriate evaluation, which lead to the significant differences in the ratio of RP+ and RP–between the training and test datasets.

The KL-6 testing is a part of the standard protocols in our hospital. Therefore, the KL-6 levels are tested for as many patients as possible. The pCT images for the training dataset were acquired between August 2003 and July 2013 using a CT scanner (Mx 8000, Philips Healthcare, Amsterdam, The Netherlands) with the following scanning parameters: tube voltage, 120 kV; in-plane pixel size, 0.98 mm; and slice thickness, 2.0 mm. Conversely, the pCT images for
the test dataset were scanned between April 2014 and March 2018 using a different scanner (Aquilion Prime, Canon Medical Systems, Otawara, Japan) with the same scanning parameters. The patient setups for the image acquisition were performed in the standardized protocol in the Kyushu University Hospital. RP grades were scored using the Common Terminology Criteria for Adverse Events version 4.0 (CTCAE v.4.0), based on clinical assessment and imaging. Anisotropic pCT images and RT structures including lung regions and gross tumor volumes (GTV) were transformed into isotropic images with an isovoxel size of 0.98 mm, using linear and shape-based interpolations, respectively.

This retrospective study was performed with the ethical approval of the institutional review board of our hospital and written informed consent was waived because of the retrospective design. All of the methods were carried out in accordance with the Declaration of Helsinki.

2.2 Development of the BN-based signature

2.2.1 Topological image processing and feature extraction. Fig 1 illustrates the overall workflow of this study. First, the most feasible set of image features (signature) was determined by evaluating their performance in the prediction of RP+. Then, the combination of the image signature and well-known predictive factors including KL-6 and dose volume indices (DVI) was explored for further improvement of the predictive performance of the model. BN-based image features were extracted from the lung regions on the axial, coronal, and sagittal planes.

![Fig 1. The overall workflow of the present study designed to achieve two major objectives: Construction of the BN-based signature and exploration of synergistic combination with known predictive factors. This study consisted of two main sections. First, the most feasible set of image features (signature) was determined by evaluating their performance in the prediction of RP+. Then, the combination of the image signature and well-known predictive factors including KL-6 and DVI was explored for further improvement of the predictive performance of the model. pCT: planning computed tomography, BN: Betti number, WD: wavelet decomposition, GLSZM: gray-level size zone matrix, GLCM: gray-level co-occurrence matrix, GLDM, gray-level dependence matrix, GLRLM: gray-level run-length matrix, NGTDM: neighborhood gray-tone difference matrix, RP: radiation-induced pneumonitis. LASSO: least absolute shrinkage and selection operator, SVM: support vector machine, KL-6: Krebs von den Lungen-6, DVI: dose volume indices.

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of the pCT images, which included the centroid of the GTV. In all, 47,616 BN-based features were obtained from 512 BN maps (256 b0 maps and 256 b1 maps) that characterize topologically invariant heterogeneous characteristics of lung regions on the pCT images by applying 93 feature calculations (18 histogram-based features and 75 texture-based features, S2 Table). The integrated histogram and texture matrices were obtained from the BN maps for the three-plane pCT images to compute the image features. The computation of the BN maps was performed by counting b0 and b1 in local regions of 256 binary images through a convolutional computation procedure using four kernel sizes (5, 7, 9, and 11 pixels) and five shifting pixels (1, 2, 3, 4, and 5 pixels) [24, 25]. To obtain the 8-bit pCT images, the original CT value range of -1350 to 150 Hounsfield units was converted to 8 bits (0 to 255). The binary images were derived from these 8-bit pCT images by thresholding, with values ranging from 0 to 255.

To compare the feasibility of the BN features with the conventional ones, 93 pCT-based features and 744 WD-based features were extracted from the original and wavelet-decomposed pCT images, respectively, using three-dimensional calculation algorithms for the same 93 features. Wavelet decomposition was performed using the high (H)-pass filter and low (L)-pass filter based on the coiflet 1 mother wavelet. Eight WD images were obtained by applying either H or L filters in the x, y, and z directions [LLL, LLH, LHL, LHH, HLL, HLH, HHL, and HHH (each of characters either L or H represents a low- or high-pass filter applied to the x, y, or z direction)].

The extraction of image features was performed using PyRadiomics package version 3.0.1 in Python 3.8.

2.2.2 Signature construction. The signatures were constructed using frequency-based voting with the LASSO-LR model. Fig 2 shows the procedure for frequency-based voting based on an imbalance adjustment strategy proposed by Schiller et al. [30]. The most frequent features were included in the signature. When features showed the same frequency (ties), those with a higher sum of absolute coefficients in the LASSO-LR models were selected. These frequencies were obtained from the non-zero coefficients of several LASSO-LR models constructed with subsets of the training dataset. The LASSO-LR model for each subset was optimized by maximizing the partial log likelihood while minimizing the L1 norm of the coefficients for the RP prediction in a leave-one-out cross-validation [31]. The subsets were constructed so that the numbers of RP+ and RP− cases were the same or similar. The RP+ cases were copied, and the RP− cases were randomly sampled to each subset. The number of subsets, p was determined using the following equation:

$$p = \left\lfloor \frac{N}{N_+} \right\rfloor,$$

where \(N_−\) and \(N_+\) indicate the number of RP− and RP+ cases, respectively. \(\lfloor \cdot \rfloor\) represents floor function. We used 11 subsets, as \(N_−\) and \(N_+\) were found to be 222 and 20, respectively (41 cases for two subsets and 40 cases for nine subsets). This strategy not only balances the impact of both groups in the construction of the LASSO-LR model but also constructs a consensus signature as a result of voting.

We also constructed a signature with a combination of the BN, pCT, and WD features (BN+pCT+WD model) to investigate the impact of the conventional image features in the RP+ prediction model. The feasibility of the conventional combination consisting of pCT and WD features (pCT+WD model) was also investigated because they were reported to perform better in combination rather than a single implementation of either pCT or WD features [13].

2.2.3 Model construction and evaluation. The occurrence of RP+ was predicted using support vector machine (SVM) models based on the signature. To balance the difference
between the number of RP+ and RP− cases, an imbalance adjustment algorithm was also applied to train the SVM models with the same subsets used in the signature construction (Fig 2). Optimal hyperparameters of the SVM were explored with respect to kernels (linear, Gaussian, and sigmoid), soft margin parameters C (ranging from 0.1 to 10), and gamma values (ranging from 0.1 to 10). Parameter optimization was performed based on the robustness index (RI), which defined higher total and lower differences in the AUCs between the training and validation. The RI between the training and validation is defined as follows:

\[
RI_{tr,va} = \frac{AUC_{tr} + AUC_{va}}{1 + |AUC_{tr} - AUC_{va}|},
\]

where \(AUC_{tr}\) and \(AUC_{va}\) are the AUCs for the training and validation, respectively. In this study, the training and validation AUCs were obtained by averaging the AUCs from the subsets (indicated by blue rhombi in Fig 2). The SVM model in each subset was validated using the leave-one-out cross-validation.

The feasibility of the models was compared with respect to the AUCs, RI, accuracy, sensitivity, and specificity based on the prediction made by ensemble averaging the outputs from reconstructed SVMs in the subsets. The reconstructed SVMs were obtained by training the
SVM model for all cases in each subset using the optimal hyperparameters. The averaged SVM output was used to calculate the AUC for the test. The number of features in the signature was explored from 1 to 10 to maximize the RI for validation and testing. RI was calculated from the AUCs for validation and testing. The accuracy, sensitivity, and specificity of the models in the test were calculated based on the ROC curves with the optimal threshold value determined by the minimization of the Euclidean distance between the ROC curve and the point representing a true positive rate of one and a false positive rate of zero.

2.3 Exploration of synergistic combination with KL-6

To assess the impact of KL-6 in combination with the BN signature, the SVM model was constructed using the combined features of the BN signature and KL-6 (BN+KL-6) based on the KL-6 dataset using the same SVM construction algorithm described in Section 2.2.3.

Since the feasibility of the DVI of lung volume receiving doses greater than 5, 10, and 20 Gy (V5, V10, and V20) and mean lung dose (MLD) has been reported in previous studies [32, 33], we also constructed an SVM model with BN and DVI (BN+DVI).

2.4 Statistical analyses

In the demographic distributions of the patients (Tables 1 and 2), the Mann–Whitney U tests were used to assess significant differences in terms of continuous quantities. In contrast, chi-squared tests were applied for the other categorical variables. Statistically significant differences and 95% confidence intervals (CIs) among the AUCs in the validation and test were assessed using the bootstrap method with 2000 iterations (significance threshold: p < 0.05). To compute the statistics regarding the AUCs, we used ensemble outputs from the reconstructed SVMs by feeding the signatures for the training and test datasets. All analyses were performed using R-4.0.3 (http://www.r-project.org/).

3. Results

Table 1 shows the demographic distribution of the patients and the significant differences between the training and test datasets. In all, the data obtained from 242 cases (20 RP+ and 222 RP–) and 30 cases (8 RP+ and 22 RP–) were included for the training and testing of the RP prediction model, respectively. Since KL-6 information was available for 218 cases out of the 272 cases, the data (KL-6 datasets) were used to explore the synergistic combination of the BN with known predictive factors. Table 2 shows the demographic distributions of the KL-6 database and the significant differences between the training and test datasets.

Fig 3 depicts circular heatmaps representing the signatures constructed by voting using the LASSO-LR models for the BN, pCT, WD, BN+pCT+WD, and pCT+WD features. Descriptions of the features are shown in S2 Table. Three, nine, and three features were selected for the signatures of BN, pCT, and WD, respectively. The signature of BN+pCT+WD consisted of only BN features. For the conventional combination of pCT+WD, the signature was composed of two features, one from the pCT and the other from the WD.

The optimal parameter for the BN maps was a kernel size of 11 and a shifting pixel of 4. The optimal SVM parameters were a sigmoid kernel, gamma value of 0.4, and C margin of 0.1 (Table 3). For the BN+pCT+WD model, the optimal parameters were a sigmoid kernel, gamma value of 0.4, and C margin of 0.1. For the pCT+WD model, the optimal parameters were a linear kernel and a C margin of 0.1. For the pCT model, the optimal parameters were a sigmoid kernel, gamma value of 2.8, and C margin of 0.1. For the WD model, the optimal parameters were a sigmoid kernel, gamma value of 1.9, and C margin of 0.1.
Table 3 summarizes the AUCs, RIs, accuracy, sensitivity, and specificity of the RP+ prediction models based on the image signatures. The AUCs and the CIs that were obtained for the validation of the BN, BN+pCT+WD, pCT+WD, pCT and WD models were 0.873 (0.777–0.968), 0.880 (0.810–0.950), 0.775 (0.681–0.867), 0.758 (0.656–0.860) and 0.785 (0.700–0.870), respectively. In the validation, BN showed a significantly higher AUC than conventional models including pCT, WD, and pCT+WD (Fig 4). The AUCs and the CIs that were obtained for the test of the BN, BN+pCT+WD, pCT+WD, pCT and WD models were 0.852 (0.697–1.000), 0.824 (0.651–0.997), 0.790 (0.620–0.959), 0.642 (0.417–0.867) and 0.545 (0.296–0.795), respectively. In the test, the BN outperformed the conventional pCT and WD models, with statistically significant differences (Fig 4). The RIs obtained for the BN, BN+pCT+WD, pCT+WD, pCT, and WD models were 1.691, 1.613, 1.541 1.255, and 1.073, respectively. In the test, the BN showed a significantly higher RI than conventional models including pCT, WD, and pCT+WD (Fig 4). The accuracies (sensitivity and specificity) of the BN, BN+pCT+WD, pCT+WD, pCT, and WD models were found to be 0.757 (0.756, 0.757), 0.724 (0.704, 0.779), 0.681 (0.664, 0.728), 0.591 (0.582, 0.616), and 0.534 (0.524, 0.562), respectively.

Table 4 summarizes the results of the exploration of synergistic combination with KL-6. The AUCs and CIs that were obtained for the validation of the BN+KL-6, BN, and BN+DVI

Table 1. Demographic distributions of the patients and significant differences between the training and test datasets.

|                                      | Training dataset | Test dataset | p-value (testing method) |
|--------------------------------------|------------------|--------------|--------------------------|
| Total number of cases                | 242              | 30           |                          |
| RP grade                             |                  |              |                          |
| Grade = 1                            | 222              | 22           | 4.952×10⁻³ (Chi-squared test) |
| Grade ≥ 2                            | 20               | 8            |                          |
| Pre-treatment KL-6 availability      |                  |              |                          |
| Available                            | 193              | 25           | 0.823 (Chi-squared test)  |
| Unavailable                          | 49               | 5            |                          |
| Pre-treatment KL-6 quantity (U/mL, min-max (median)) | 115–1853 (262) | 134–678 (283) | 0.745 (Mann-Whitney U-test) |
| Age (y, min-max (median))            | 51–92 (77)       | 54–90 (74)   | 0.722 (Mann-Whitney U-test) |
| Sex                                  |                  |              |                          |
| Male                                 | 150              | 21           | 0.511 (Chi-squared test)  |
| Female                               | 92               | 9            |                          |
| Stage                                |                  |              | 0.146 (Chi-squared test)  |
| T2N0M0 (stage IB)                    | 185              | 27           |                          |
| Dose and fraction                    |                  |              |                          |
| 12 Gy × 4 Fr. (48 Gy)                | 236              | 25           | 4.00 × 10⁻² (Chi-squared test) |
| 13 Gy × 4 Fr. (52 Gy)                | 2                | 3            |                          |
| 6 Gy × 10 Fr. (60 Gy)                | 4                | 2            |                          |
| Dose prescription method             |                  |              |                          |
| Isocenter                            | 142              | 10           | 1.200 × 10⁻² (Chi-squared test) |
| D95 for PTV                          | 100              | 20           |                          |
| Pre-existing pulmonary diseases      |                  |              |                          |
| IP                                   | 7                | 1            | 0.196 (Chi-squared test)  |
| COPD                                 | 35               | 8            |                          |
| IP and COPD                          | 3                | 1            |                          |
| Neither                              | 197              | 20           |                          |

RP: radiation pneumonitis, KL-6: Serum Krebs von den Lungen-6, PTV: planning target volume, IP: interstitial pneumonia, COPD: chronic obstructive pulmonary disease.

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models were 0.901 (0.818–0.985), 0.873 (0.777–0.968), and 0.905 (0.819–0.992), respectively. The AUCs and CIs that were obtained for the test of the BN+KL-6, BN, and BN+DVI models were 0.825 (0.647–1.000), 0.807 (0.633–0.999), and 0.684 (0.434–0.587), respectively. BN+KL-6 showed higher AUCs than the BN model for both the validation and test. In contrast, although the BN+DVI model showed a higher AUC for the validation, it showed a significantly lower AUC than the BN model for the test (Fig 4). The RIs obtained for the BN+KL-6, BN, and BN+DVI models were 1.603, 1.576, and 1.302, respectively. The accuracies (sensitivity and specificity) of the BN+KL-6, BN, and BN+DVI models were found to be 0.724 (0.711, 0.765), 0.708 (0.699, 0.738), and 0.621 (0.618, 0.634), respectively.

4. Discussion
The results of the feature selection suggested the robustness and superiority of the BN features for RP+ prediction. The conventional pCT and WD features were not selected in the signature, although they were also included as candidates for the signature. The existence of conventional features affected the course of the feature selection based on voting, because the signature based on the BN+pCT+WD included nine features composed of six additional features apart from the three features selected in the BN-based signature. Nevertheless, the AUCs of the BN and BN+pCT+WD models did not show statistically significant differences in either the
Topological imaging signature and biomarker synergy for radiation pneumonitis prediction

**Feature type**
- Betti number (BN)
- Planning computed tomography (pCT)
- Wavelet decomposition (WD)

**Z-score**

**Radiation induced pneumonitis (RP) status**
- RP –
- RP +
validation or test (Fig 4). Even though the constituents for the signature changed, the representative BN features were selected under consensus by voting with multiple LASSO-LR models, which consistently produced higher AUCs than the conventional models.

Table 5 compares the AUCs that were obtained in the present study to the AUCs obtained in previous studies. Our BN-based model outperformed the models reported in previous studies for RP prediction in SABR for lung cancer with respect to the test AUCs and RI [12, 13]. Although Hirose et al. reported a higher AUC for the validation than the BN model using the pCT+WD features, their model showed a lower AUC for the test [13]. In our study, we attempted to balance the AUCs for the pCT+WD model for the validation and test using RI. However, it showed a similar performance to the one reported by them. These results imply that the feasibility of conventional pCT and WD features is quite limited in RP prediction before SABR for lung cancer.

The BN+KL-6 model improved the prediction, although it did not show a significant difference in comparison with the BN model. Although the use of KL-6 only cannot provide sufficient predictive performance, it could support RP+ prediction using the BN signature. All evaluation criteria including the RI, AUCs, accuracy, sensitivity, and specificity were improved in the BN+KL-6 model (Table 4). Although further investigations using larger datasets are needed, they may offer a reliable system for evaluating RP risk in lung cancer patients.

We also constructed the SVM model with BN-based signature, KL-6, and the status of prePDs. However, the model showed a similar predictive performance with the BN+KL-6 model. The AUC values for the validation and the test were 0.872 and 0.816, respectively. This might be because the model included the important phenotypes of the IP or COPD which is associated with the occurrence of the RP. The distributions of the outputs from the SVM model based on the BN+KL-6 showed a correlation with the status of pre-existing lung diseases (Table 6). Most of the IP-positive cases were predicted as RP+ by the model. On the other hand, two-third of all the COPD-positive cases were predicted as RP–. Those distributions of prePDs are similar to the observation reported in the previous studies; higher RP+ risk for the IP and lower RP+ risk for the COPD [14–18].

Table 3. Summary of the areas under the receiver operating characteristics curves (AUCs) and robustness indices (RIs) obtained between the validation and test for the image signatures.

| Number of features in the signature | Optimal SVM parameters | AUC (95% CI) | RI | Accuracy | Sensitivity | Specificity |
|-----------------------------------|-------------------------|--------------|----|----------|-------------|-------------|
|                                   | Kernel | Gamma | C margin | Validation | Test |                |              |              |
| BN                                | 3      | Sigmoid | 0.4 | 0.1 | 0.873 (0.777–0.968) | 0.852 (0.697–1.000) | 1.691 | 0.757 | 0.756 | 0.757 |
| BN+pCT +WD                        | 8 (all BN) | Sigmoid | 0.4 | 0.1 | 0.880 (0.810–0.950) | 0.824 (0.651–0.997) | 1.613 | 0.724 | 0.704 | 0.779 |
| pCT+WD                            | 2 (1 pCT, 1 WD) | Linear | - | 0.1 | 0.775 (0.681–0.867) | 0.790 (0.620–0.959) | 1.541 | 0.681 | 0.664 | 0.728 |
| pCT                                | 9      | Sigmoid | 2.8 | 0.1 | 0.758 (0.656–0.860) | 0.642 (0.417–0.867) | 1.255 | 0.591 | 0.582 | 0.616 |
| WD                                | 3      | Sigmoid | 1.9 | 0.1 | 0.785 (0.700–0.870) | 0.545 (0.296–0.795) | 1.073 | 0.534 | 0.524 | 0.562 |

BN: Betti number, pCT: Planning computed tomography, WD: Wavelet-decomposition.

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We investigated Spearman’s correlation coefficients between KL-6 and the three features in the BN-based signature and found no correlations among them (correlation coefficient < 0.17). Therefore, the BN-based signature and KL-6 provide independent and useful measurements for RP+ prediction based on lung tissue conditions.

Table 4. Summary of the areas under receiver operating characteristics curves (AUCs) and robustness indices (RIs) obtained between the validation and test for the Betti number (BN)-based signatures with serum Krebs von den Lungen-6 (KL-6) or dose volume indices (DVI).

| Number of features in the signature | Optimal SVM parameters | AUC (95% CI) | RI | Accuracy | Sensitivity | Specificity |
|------------------------------------|------------------------|--------------|----|----------|-------------|-------------|
| BN+KL-6 4 (3BN, 1KL-6)             | Sigmoid, 0.1, 0.4      | 0.901 (0.818–0.985) | 0.825 (0.647–1.000) | 1.603 | 0.724 | 0.711 | 0.765 |
| BN 3                                 | Sigmoid, 0.4, 0.1      | 0.873 (0.777–0.968) | 0.807 (0.633–0.999) | 1.576 | 0.708 | 0.699 | 0.738 |
| BN+DVI 7 (3BN, 4DVI)               | Sigmoid, 0.1, 0.4      | 0.905 (0.819–0.992) | 0.684 (0.434–0.587) | 1.302 | 0.621 | 0.618 | 0.634 |

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The association between KL-6 levels and the severity of IP has been reported in previous studies [4, 34]. In this study, we found that KL-6 could detect patients with and without pre-existing IP in the training, as shown in the top right and bottom left examples in Fig 5, respectively. However, an increase in KL-6 does not always indicate the presence of IP. Yoshimatsu et al. reported that KL-6 levels could increase by the existence of lung cancer itself, with a significant association with tumor size [10]. Several cases showed high pre-treatment KL-6 levels, although they did not have pre-existing IP (the bottom right example in Fig 5). The sensitivity and specificity for the prediction of RP with KL-6 were found to be 0.582 and 0.625, respectively, using the optimal threshold value of 282.5 U/mL (vertical blue line in Fig 5), which is quite similar to that reported by Iwata et al. [9]. Contrastingly, the BN+KL-6 model could successfully predict the RP status in these patients. Furthermore, the model could predict the occurrence of RP in cases without IP or COPD (top left example in Fig 5). The sensitivity and specificity for the RP+ prediction with the BN+KL-6 model were found to be 0.746 and 0.938, respectively, using the optimal threshold value of 0.383 (horizontal blue line in Fig 5).

Variability of dose prescription methods and dose fractions may cause the lower performance of the BN+DVI model. In our database, which we have summarized in Tables 1 and 2, SABR plans were constructed using two dose prescription methods (isocenter or D95 for planning target volume) and three dose fractions [12 Gy×4 Fr. (48 Gy), 13 Gy×4 Fr. (52 Gy) or 6 Gy×10 Fr. Gy (60 Gy)]. Furthermore, there were statistically significant differences in these parameters between the training and test datasets. These differences affect the DVIs and can be attributed to poor prediction results with the BN+DVI model in the test.

The present study had four limitations. First, the BN model could not entirely evaluate the three-dimensional textures of the lung regions. Although we extended the range of the computation to the z-direction using the three cross-sections, it still utilizes two-dimensional computation algorithms for b0 and b1. Two-dimensional BNs cannot distinguish holes as intersections of the trachea from cavitation due to IP or COPD. Second, our model did not consider the impact of X-ray irradiation in SABR. As mentioned above, the DVIs did not improve the performance of RP+ prediction in our datasets. However, with respect to the RP

Table 5. Comparison of areas under the receiver operating characteristics curves (AUCs) obtained in the present study to the AUCs obtained in previous studies.

| Number of cases | Dataset | Feature type | AUC Validation | AUC Test | RI |
|-----------------|---------|--------------|---------------|---------|----|
| The present study | 218 | Training 193 BN+KL-6 | 0.901 | 0.825 | 1.603 |
| | Testing 25 BN | 0.873 | 0.807 | 1.576 |
| Hirose et al | 275 | Training 245 pCT+WD | 0.871 | 0.756 | 1.459 |
| | Testing 30 pCT | 0.750 (not validated) | – |
| Morgan et al | 14 | Not validated pCT | 0.750 (not validated) | – |

BN: Betti number, KL-6: Serum Krebs von den Lungen-6, pCT: planning computed tomography, WD: wavelet decomposition.

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Table 6. Relationships between the predictions using the BN+KL-6 model and the pre-existing pulmonary diseases in the training dataset.

| SVM prediction (BN+KL-6) | Status of pre-existing pulmonary diseases | Total |
|--------------------------|-----------------------------------------|-------|
| IP                       | COPD                                    | Neither |
| RP+                      | 6                                       | 42    | 60 |
|                          | 11                                      | 2     | 133 |
| RP–                      | 1                                       | 2     | 110 |

BN: Betti number, KL-6: Serum Krebs von den Lungen-6, IP: interstitial pneumonia, COPD: chronic obstructive pulmonary disease.

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+ prediction based on dosimetric information, Adachi et al. developed a model that implemented dosimetric features [35]. They showed a remarkable RP+ prediction performance that was much better than the DVI-based prediction. The dosimetric features could provide a critical dosimetric factor for RP+ prediction. Third, the present study could not clarify the feasibility of the model against other types of irradiation. Previous studies reported that RP could occur in several types of radiation treatments, such as conventional fractionated radiotherapy for lung cancer [36, 37] or head and neck regions [38]. Our model may reflect the vulnerability of lung regions to SABR for lung cancer. However, the generalizability of the findings obtained in the present study should be investigated in a wide variety of radiation treatments to provide comprehensive support for the risk assessment of RP in clinical practice. Last, we have not performed standardizations to normalize the impacts derived from the variations in the CT scanners. Since the BN features showed their robustness against the variations in the CT scanners in our previous study [25], we assumed that the BN features could mitigate the differences of the scanners in the training and testing datasets. However, it could show better prediction performance when the standardization was performed before the analysis. Haga et al reported that the classification performance of the histological subtypes of the early-stage lung cancer based on the imaging features extracted from CT images could be improved when standardization methods such as min-max normalization, z-score, or whitening from the principal component analysis were applied to the different (training and test) databases with different

Fig 5. Distributions of the pretreatment serum Krebs von den Lungen-6 (KL-6) and support vector machine (SVM) outputs based on BN +KL-6. Representative features and Betti number (BN, b0) maps were shown for the cases where the SVM model could successfully predict the radiation pneumonitis (RP) status [positive (+) or negative (−)]. Labels of the BN maps (b0) are followed by the threshold value for obtaining the BN maps. Hist: histogram, NGTDM: neighborhood gray-tone difference matrix, GLSZM: gray-level size zone matrix.

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scanning conditions [39]. We intend to investigate the impact of standardization on the classification performance of the BN features in future work.

5. Conclusions

This study demonstrated the comprehensive performance of BNs for the prediction of potential RP+ patients before SABR for lung cancer. This prediction could be enhanced in combination with pretreatment with KL-6. We considered that the BN+KL-6 model was more feasible for the RP+ prediction because it showed the highest AUC value in the test of the exploration of synergistic combinations. These predictions could help radiation oncologists in decision making for dose reduction to avoid severe RP+ in personalized SABR or selection of alternative treatments to provide a better quality of life for each patient. Further studies with larger datasets and different radiation treatments to support the risk assessment of RP in clinical practice are needed.

Supporting information

S1 Table. Prediction performances of serum Krebs von den Lungen-6 (KL-6) and dose volume indices (DVI) of lung volume in the dataset used in the present study. The support vector machines were constructed for KL-6 and DVIs using the same ensemble strategy described in the main article. (DOCX)

S2 Table. Radiomic features with the feature types. (DOCX)

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