Research paper

Tumor grading of soft tissue sarcomas using MRI-based radiomics

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A B S T R A C T

Background: Treatment decisions for multimodal therapy in soft tissue sarcoma (STS) patients greatly depend on the differentiation between low-grade and high-grade tumors. We developed MRI-based radiomics grading models for the differentiation between low-grade (G1) and high-grade (G2/G3) STS.

Methods: The study was registered at ClinicalTrials.gov (number NCT03798795). Contrast-enhanced T1-weighted fat saturated (T1FSGd), fat-saturated T2-weighted (T2FS) MRI sequences, and tumor grading following the French Federation of Cancer Centers Sarcoma Group obtained from pre-therapeutic biopsies were gathered from two independent retrospective patient cohorts. Volumes of interest were manually segmented. After preprocessing, 1394 radiomics features were extracted from each sequence. Features unstable in 21 independent multiple-segmentations were excluded. Least absolute shrinkage and selection operator models were developed using nested cross-validation on a training patient cohort (122 patients). The influence of CombBatHarmonization was assessed for correction of batch effects.

Findings: Three radiomic models based on T2FS, T1FSGd and a combined model achieved predictive performances with an area under the receiver operator characteristic curve (AUC) of 0.78, 0.69, and 0.76 on the independent validation set (103 patients), respectively. The T2FS-based model showed the best reproducibility. The radiomics model involving T1FSGd-based features achieved significant patient stratification. Combining the T2FS radiomic model into a nomogram with clinical staging improved prognostic performance and the clinical net benefit above clinical staging alone.

Interpretation: MRI-based radiomics tumor grading models effectively classify low-grade and high-grade soft tissue sarcomas.

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1. Introduction

Soft tissue sarcomas (STS) constitute rare malignant diseases [1]. In contrast to many other malignant tumors, treatment decisions strongly depend on pre-therapeutic tumor grading [2].

Tumor grading is commonly determined during histological workup of pre-therapeutic biopsies. Two distinct grading systems were developed by the National Cancer Institute and the French Federation of Cancer Centers Sarcoma Group (FNCLCC) [3,4]. In direct comparison, the FNCLCC system appeared to predict distant metastasis development.
Research in context
Evidence before this study

Therapy decisions for patients with soft tissue sarcomas (STS) vastly depend on the differentiation between low-grade and high-grade STS. In contrast to low-grade STS, patients diagnosed with high-grade STS have an unfavorable prognosis and often suffer from the occurrence of distant metastases. As consequence, patients with high-grade STS receive multimodal therapy regimens including radiation therapy and/or chemotherapy. So far, tumor grading is defined by histological work up after invasive biopsies. Previous studies, could identify semantic magnetic resonance imaging (MRI) derived properties, such as contrast enhancement, necrosis or tumor heterogeneity, associated with higher tumor grading. First small single center studies were able to demonstrate an association of quantitative radiomics features with tumor grading, too.

Added value of this study

We have developed non-invasive tumor grading models for the differentiation between low-grade and high-grade sarcomas using MRI-based radiomics. All models were validated using an independent external patient cohort. The radiomic classifier based on contrast enhanced T1-weighted MRI sequences achieved significant patient risk stratification. Combining the T2-weighted sequence based radiomic model into a nomogram with clinical stages improved the clinical net benefit and prognostic performance above clinical staging alone.

Implications of all the available evidence

The study provides radiomic MRI-based models to non-invasively differentiate low-grade from high-grade sarcomas. The presented models may be used as a non-invasive grading estimate if the pathological workup does not yield a clear result, or the tumor exhibits inhomogeneous areas that are difficult to access via CT-targeted or open biopsy. Moreover, the proposed nomogram may be used for improved prognostic assessment prior to therapy.

2. Methods

2.1. Patients

The trial was registered at ClinicalTrials.gov (NCT03798795). All patients with histologically proven STS with known FNCLCC tumor grading determined by biopsy prior to therapy and availability of contrast-enhanced T1-weighted fat saturated (T1FSGd) as well as fat-saturated T2-weighted (T2FS) MRI sequences of pre-therapeutic MRI were included. Patient records were analyzed for patient demographics (see Table 1) and histological subtypes (see Supplemental Table 1) from two independent retrospective patient cohorts from the Technical University of Munich (TUM) (138 patients, one patient with two...
independent STS) and the University of Washington, Seattle (UW) (139 patients). Exclusion criteria were previous RT, Ewing sarcoma, primary bone sarcomas, and endoprosthesis-dependent MRI artifacts. Approval from the ethics committee was received in both institutions. Patients were treated after informed consent. Overall survival (OS) was calculated from the initial pathologic diagnosis to the time point of death or the time point of censoring.

2.2. Image acquisition and definition of volume of interests

Complete imaging studies (availability of both sequences of interest) were found in 122 patient in the TUM cohort and 103 patients in the UW cohort (see patient workflow in Supplemental Fig. 1). See Supplemental Tables 2 and 3 for MRI vendors and acquisition parameters.

Tumor segmentation was conducted manually using MIM software version 6.6 at UW by MS, MM and TC (MIM Software Inc., Cleveland, USA), Eclipse 13.0 (Varian Medical Systems, Palo Alto, USA), and iplan RT 4.1.2 (Brainlab, Munich, Germany) at TUM by JP. The volume of interest (VOI) was defined as the primary tumor excluding surrounding edematous changes. Multiple delineations were performed for 21 randomly selected patients by three radiation oncology residents (JP, TA, MS) in the TUM cohort (see Fig. 1) to compensate for operator-dependent bias. For comparison, dice coefficients (DC) were calculated using the DiceComputation module of 3D Slicer (3D Slicer, Version 4.8 stable release) [19].

2.3. Image preprocessing and radiomics feature extraction

Feature extraction, model building and statistical analyses were performed by JP and AO. In order to compensate for non-uniform intensity caused by field inhomogeneity, N4ITK MRI bias field correction was applied to each imaging study using Slicer3D implementation (3D Slicer, Version 4.8 stable release) [20]. The pyradiomics (version 2.1) implementation in python (version 3.6.4) was utilized for further preprocessing steps and radiomics feature extraction [21]. Intensity normalization redistributed the image at the mean with the standard deviation and a scale of 100. A fixed bin width of 10 was used for image discretization (a detailed description is provided in the supplemental material). Isotropic resampling to a voxel size of 3 × 3 × 3 mm was performed using B-spline interpolation. All models were also calculated using a voxel size of 1 × 1 × 1 mm showing inferior predictive performances.

Image reconstruction was performed applying wavelet decompositions filtering and Laplacian of Gaussian filtering with sigma values of 3.0, 4.0, and 6.0. Further on, local binary pattern-derived images were calculated using a level of one and two as well as the kurtosis image. 1394 features were extracted from filtered and original images in three dimensions including shape features, first-order features, and texture features. All extracted features are listed in Supplemental Table 4.

2.4. Feature reduction and predictive model building

Feature reduction, batch correction, modeling, and statistical analyses were performed in R (version 3.4.0, R core team, Vienna, Austria). All features susceptible to minor segmentation variances in the 21 patients that received multiple independent segmentations (intraclass correlation coefficient (3,1) of <0.8) were excluded.

For model building and unbiased performance evaluation on the training set, 10 iterations of five-fold nested cross validation were performed using the code published by Deist et al. built upon the training set, 10 iterations of correlation coefficients that received multiple independent segmentations (intraclass variance). All features susceptible to minor segmentation variances in the 21 patients were then split into five subgroups (inner folds). Each Subgroup was then split once more for five times (outer folds). Hyperparameters were optimized as part of the inner folds. The selected hyperparameters were then used for testing on the five outer folds. The total mean area under the receiver operator characteristic (ROC) curve (AUC) over all outer folds was calculated for model comparison (see Supplemental Fig. 2 for a graphical depiction). The hyperparameter combination with the best mean performance was used to retrain a final model on the whole training set. Model performance with the optimal hyperparameter combination was assessed using cross validation as described above.

First, we compared the performance of seven machine learning techniques with inbuilt feature reduction (elastic net logistic regression, least absolute shrinkage and selection operator (LASSO), random forest, LogitBoost, decision tree, support vector machine and neural network). The influence of additional feature reduction by principal component analysis (PCA), resampling and reweighting were tested (see Supplemental Table 6) [22]. None of the techniques improved the performance further. We decided to apply the LASSO method without prior PCA as it showed good performance in cross validation, model simplicity and low tendency to overfitting (see supplemental material for method description and Supplemental Table 7 for selected hyperparameter values). Previous studies have demonstrated a competitive performance compared to other machine learning methods [22,23]. At the same time, it allows for a better interpretability due to the direct linear relationship between input features and outcome.

All models were trained to classify high-grade STS (G1) vs. low-grade STS (G2/G3). Three radiomic models Radiomics-T1FSGd, Radiomic T2FS, and Radiomics-combined were developed using the respective features as input. For comparison, logistic regression models of Tumor-
Volume alone, a Clinical model based upon “Age” and the TNM-staging system, and a combined model Clinical-Volume-combined were evaluated.

Finally, we trained elastic net regression models for the prediction of overall survival (OS) using five-fold cross validation using the “glmUtils” package. Model performance of all final models was finally determined on the external validation patient cohort (UW) to get an unbiased result.

2.5. Batch correction via ComBatHarmonization

Recently the ComBatHarmonization has been proposed as a method for the correction of batch effects among radiomics multicenter cohorts [24]. Its value to potentially improve reproducibility between different centers has been shown in multiple studies [25,26]. Based on the given feature distribution it estimates the additive and multiplicative batch effects using a maximum likelihood approach. We applied the R ComBat script (https://github.com/jfortin1/ComBatHarmonization) correcting for MRI scanner models [27]. Further on, we evaluated the effect of using STS histology as a biological covariate.

2.6. Decision curve analysis

In order to analyze the clinical usefulness of the developed classifiers, decision curve analysis was conducted as described by Vickers et al. [https://www.mskcc.org/departments/epidemiology-biostatistics/biostatistics/decision-curve-analysis] [28]. The net benefit is calculated by subtracting the proportion of false-positive patients from the proportion of true-positive patients, weighted by the relative harm of a false-negative and false-positive result. As a reference, the decision curves for treating all patients and treating no patients are displayed. A decision models shows a clinical benefit if it achieves larger net benefit values than both reference strategies.

2.7. Statistical analysis

ROC curves and respective AUC values were calculated and compared using the 1000-fold bootstrapping as implemented in the “fbrroc” and “pROC” packages [29]. Patient stratification for survival was tested via Kaplan-Meier survival analysis using the “ggkm” package. Dichotomization was performed using the class output provided by the glmnet function inside of the “caret” package. Statistical significance was assessed using the log-rank test. Univariate logistic regression was used to assess correlation to tumor grading using the “MCC” computed on the “Gray level co-occurrence matrix” (GLCM) and “DependenceEntropy” derived from the “Gray Level Dependence Matrix” (GLDM) (wavelet.HLH_gldm_MCC and wavelet.HLH_gldm_DependenceEntropy) were among the three most important features.

With AUC-values of 0.77 and 0.78 the models Radiomics-T2FS and Radiomics-T1FSGd showed a similar predictive performance (p = .427) on the training set (see Table 2 for AUC values and 95% confidence intervals [95CI]). The Radiomics-combined model achieved a significantly better predictive capacity (AUC 0.84) than Radiomics-T2FS and Radiomics-T1FSGd (p < .001, respectively). The Clinical model (AUC: 0.55) and the Tumor-Volume model (AUC: 0.44) did not show a predictive capacity better than random. Combining clinical parameters and tumor volume as “Clinical-Volume-combined” model did not increase predictive performance further (AUC: 0.52). All three models performed significantly worse than all radiomic model (p < .001, for each comparison).

3.2. Training of the radiomic classifiers for tumor grading prediction

After radiomics feature reduction due to segmentation variability 1309, 1340, and 2609 features were used for the development of the Radiomics-T1FSGd, Radiomics-T2FS and Radiomics-combined models, respectively. During training of Radiomics-T2FS, Radiomics-T1FSGd, and Radiomics-combined, 24, 22 and 13 features with non-zero coefficients were selected, respectively (see Supplemental Table 8 for selected features and feature importance ranking). Features were selected from various feature types and filtering methods. The most important T2FS-based features were first order intensity features based on wavelet filtering (wavelet.HLH_firstorder_Kurtosis and wavelet.HLH_firstorder_Mean), and the local binary pattern texture featurebusy (lbp.3D. k_rgttdm_Busyness). The T1FSGd based model was similarly dominated by wavelet.HLH_firstorder_Mean as for T2FS. In addition, wavelet decomposition based texture features such as “Maximum Correlation Coefficient” (MCC) computed on the “Gray level co-occurrence matrix” (GLCM) and “DependenceEntropy” derived from the “Gray Level Dependence Matrix” (GLDM) (wavelet.HLH_gldm_MCC and wavelet.HLH_gldm_DependenceEntropy) were among the three most important features.

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3.3. Independent validation of the developed classifiers

The predictive models were evaluated on the validation set after batch correction. The T2FS-based model showed good reproducibility with an AUC of 0.78 (absolute difference of +0.01) which was significantly better than Clinical but not Tumor-Volume (p = .01 and p = .056, respectively) (see Table 2 for AUC values and 95%CI, Fig. 2 for ROC curves, and Supplemental Fig. 3 for calibration curves). Validation of the Radiomics-T1FSGd model showed a larger drop in predictive performance with an absolute difference of −0.09 and an AUC of 0.69. The Radiomics-combined model showed a similar performance than the T2FS model (AUC 0.76, absolute difference −0.09) which was, however, not significantly better than Clinical or Tumor-Volume (p = .056 and p = .165, respectively). The observed difference in between the radiomic model did not reach statistical significance (Radiomics-T1FSGd vs. Radiomics-T2FS, p = .192, Radiomics-T1FSGd vs. Radiomics-combined, p = .348). The Clinical model and the Tumor-Volume model showed no predictive performance better than random. Clinical-Volume-combined, however, showed predictive performance significantly better than random with an AUC of 0.67.

In contrast to Clinical, Tumor-Volume, Clinical-Volume-combined, and Radiomics-T1FSGd, Radiomics-T2FS and Radiomics-combined were significantly associated with tumor grading in univariate logistic regression analysis in the independent validation cohort (odds ratio (OR) for a 10% increase in predicted probability: Clinical OR = 1.4 (95%CI 0.62-3.3) p = 1.0, Tumor-Volume OR = 1.5 (95%CI 0.9-6.1) p = .33, Radiomics-T1FSGd OR = 1.4 (95%CI 1.09-1.86) p = .064, Radiomics-
The Radiomics-combined model showed the highest accuracy (0.83) and sensitivity (0.90), but low specificity (0.50), for the prediction of high-grade STS compared to Radiomics-T2FS (accuracy: 0.78, sensitivity: 0.87, specificity: 0.40) and Radiomics-T1FSGd (accuracy: 0.78, sensitivity: 0.87, specificity: 0.40) (see Supplemental Table 9 for additional performance metrics) on the validation set.

3.4. Influence of batch correction

The influence of ComBatHarmonization on validation performance was different depending on the predictive model. The T1FSGd-based models Radiomics-T1FSGd improved with an absolute AUC difference of +0.08. Radiomics-T2FS and Radiomics_combined showed less marked increases in AUC of +0.03 and +0.04, respectively. Application of histology as biological covariate did not increase predictive performance

### Table 2

Predictive performance metrics of the radiomic classifiers.

| Model                        | AUC   |
|------------------------------|-------|
| Clinical-Volume-combined     | 0.52 (0.49-0.55) | 0.67 (0.52-0.8) |
| Clinical                     | 0.44 (0.41-0.46) | 0.57 (0.43-0.7) |
| Tumor-Volume                 | 0.55 (0.52-0.58) | 0.62 (0.47-0.75) |
| Radiomics-T1FSGd             | 0.78 (0.75-0.81) | 0.69 (0.57-0.81) |
| Radiomics-combined           | 0.84 (0.81-0.86) | 0.76 (0.62-0.88) |
| Radiomics-T2FS               | 0.77 (0.75-0.79) | 0.78 (0.66-0.89) |

Area under the receiver operator characteristic curve (AUC) values for the differentiation of high-grade from low-grade soft tissue sarcomas. 95% confidence intervals are shown in parentheses.

**Fig. 2.** Predictive performance of radiomics tumor grading models. Receiver operator characteristic curves (ROC) and the respective area under the curve (AUC) values depicting the performance of the prediction models Clinical, Clinical-Volume-combined, Tumor-Volume, Radiomics-T1FSGd, Radiomics-T2FS, and the Radiomics-combined on the validation cohort. The shaded blue areas depict the 95% confidence interval which is shown in parentheses. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
profundely rising the performance of Radiomics-T1FSGd to 0.70 while decreasing performances of Radiomics-T2FS and Radiomics-combined (AUC of 0.75 and 0.76, respectively) (see Supplemental Table 10).

3.5. Patient risk stratification

Tumor grading (high-grade vs. low-grade) significantly stratified patients for OS in the training and validation patient cohorts (p = 0.0006 and p = 0.045, respectively) (see Fig. 3 for Kaplan Meier survival curves and p-values). We used the classification of the developed radiomics grading models for dichotomization of the patient cohort into low-risk and high-risk patients. On the training set, all models achieved a separation of survival curves. The difference in survival was significantly different for the models Radiomics-T1FSGd (p = .0318), Radiomics-T2FS (p = .00328), and Radiomics-combined (p = .0204) but not for Clinical-Volume (p = .334). On the validation set, Radiomics-T1FSGd achieved a separation of survival curves with significant patient stratification (p = .0268). Interestingly, there was no significant separation of survival curves for the three other models.

3.6. Development of a clinical radiomics nomogram

Next, we analyzed if the proposed models may provide an incremental benefit above the existing clinical staging system. We generated multivariate nomograms by combining the AJCC staging system (7th edition) with the respective grading models. The model based on Radiomics-T2FS showed the best predictive performance for OS in the validation set with a concordance index (C-index) of 0.74 (95%CI 0.64–0.84) in comparison to Radiomics-T1FSGd (C-index: 0.71, 95%CI 0.61–0.81), Radiomics-combined (C-index: 0.72, 95%CI 0.62–0.83), and to the AJCC staging system alone (C-index: 0.69, 95%CI 0.60–0.78). For comparison, tumor volume alone and inside of a similar multivariate model achieved a C-index of 0.54 (95%CI 0.43–0.65) and 0.71 (95%CI 0.61–0.81), respectively. See Supplemental Table 11 for all training and testing performances.

For the best model based on Radiomics-T2FS, a nomogram was created (see Fig. 4a). With an AUC of 0.82 for 2-year survival, the model showed good discrimination and good calibration (Fig. 4b and c). Further on, it achieved significant patient risk stratification (p < .0001) (Fig. 4d). In decision curve analysis, the Radiomics-T2FS nomogram outperformed the “treat all” and “treat none” strategies, as well as the AJCC staging system and tumor grading alone in the threshold probability range between 0.2 and 0.55.

3.7. Prognostic radiomic prediction models show moderate performance

Finally, we trained radiomic prediction models directly for OS. The models based on the T1FSGd features, T2FS features, the combined feature set, and Tumor-Volume achieved predictive performances with C-indices of 0.55 (95%CI 0.45–0.65), 0.60 (95%CI 0.49–0.70), 0.60 (95%CI 0.50–0.69), and 0.54 (95%CI 0.43–0.65), respectively. Combining the AJCC staging system and the above mentioned models did not confer an incremental benefit (C-indices for T1FSGd: 0.67 (95%CI 0.57–0.77), T2FS: 0.70 (95%CI 0.59–0.80), combined features: 0.64 (95%CI 0.55–0.74), and Tumor-Volume: 0.71 (95%CI 0.61–0.81)).

4. Discussion

We have developed MRI-based radiomic models for tumor grading of STS focusing on the differentiation of high-grade STS from low-grade STS. A T2FS-sequence-based model showed a good reproducibility with the best performance on the independent validation patient set. A radiomic model involving T1FSGd-based features demonstrated a larger drop in predictive performance which was partly improved by ComBat harmonization. The Radiomics-T1FSGd model achieved significant patient risk stratification for OS. A nomogram combining Radiomics-T2FS and the AJCC clinical staging system achieved the best prognostic performance with significant patient stratification and a larger clinical net benefit than AJCC staging system alone. Radiomic models directly trained for the prediction of OS showed only moderate performances.

Previous analyses demonstrated a correlation of qualitative MRI features such as peritumoral contrast enhancement with tumor grading [30]. In a recent small retrospective study encompassing 19 patients, diffusion weight MRI-based radiomics features were used to distinguish G2 and G3 STS [31]. A further study used T2FS sequence based radiomics to discriminate low-grade from high-grade STS in a small pilot study of 35 patients [32]. Although patient numbers were substantially low and
no external validation cohorts were used, the studies demonstrated the principal potential to correlate quantitative radiomic features to tumor grading. A further study could demonstrate the differentiation of G2 and G3 STS using computer tomography-based radiomics albeit with a low predictive performance with an AUC of 0.65 [18]. Although radiomics features seem to provide information regarding the differentiation of G2 or G3 STS, there is currently no clinical benefit for such a differentiation. Therefore, we focused on the clinically relevant differentiation of high-grade from low-grade STS.

An imaging-based prognostic classifier as developed in this study could be applied clinically in multiple scenarios. First, in the rare case that a biopsy is clinically difficult to perform due to the anatomical location, the radiomic classifier could be used instead to identify high-grade tumors as a basis for therapy decisions. Further practical applications may involve tumors that exhibit inhomogeneous areas that are difficult to access or have different access routes in CT-targeted or open biopsies. Secondly, if the pathological work-up does not yield a clear result the proposed models may be used as an additional biomarker. Thirdly, in radiotherapy planning the spatial distribution of relevant radiomics features could be used as a basis for dose painting following the radiomics target volume concept [9,33]. For instance, radiation dose could be escalated in high-grade subvolumes. Finally, as proposed in this study the MRI radiomic grading model may be combined with clinical staging providing complementary information and improving prognostic stratification.

In the current study we restricted the volumes of interest to the gross tumor volume excluding surrounding edematous changes in both sequences. By this, we hoped to ensure a better comparability of T2FS and T1FSGd based models. Secondly, inclusion of edematous changes may produce less reproducible segmentations and radiomic feature values could be influenced by the kind of the surrounding tissue. On the contrary, by using this approach we may have missed additional information such as measures for tissue infiltration.

Fig. 4. A clinical radiomics nomogram. A multivariate nomogram combining the Radiomics-T2FS prediction model with the AJCC clinical staging system is illustrated (a). The receiver operator characteristic (ROC) curve and the representative area under the curve (AUC), as well as the calibration curve, each at two years, are shown (b,c). The Kaplan Meier survival curve for patients’ overall survival displaying patient stratification by the proposed nomogram is depicted (d). Finally, decision curve analysis was performed comparing the net benefit by the Radiomics-T2FS nomogram with the AJCC clinical stage and tumor grading alone. The net benefit is calculated by subtracting the proportion of false-positive patients from the proportion of true-positive patients, weighted by the relative harm of a false-negative and false-positive result [28]. The threshold probability was calculated for death after five years. For reference, the two strategies “treat all” and “treat none” are displayed. A decision model shows a clinical benefit if the respective curve shows larger net benefit values than both reference strategies.
Further on, we analyzed the helpfulness of radiomic models for prognostic assessment. Radiomic models directly trained to predict OS only showed moderate predictive performances. The Radiomics-T1FSGd grading model showed the propensity of significant patient risk stratification in the validation set in contrast to the models with the highest performance in predicting tumor grade (Radiomics-T2FS and Radiomics-combined). On the contrary, Radiomics-T1FSGd showed less improvement in prognostic performance in combination with the AJCC staging system than Radiomics-T2FS and Radiomics-combined. Considering that the T1FSGd-based model trained directly for OS showed worse predictive capacity than T2FS-based models contradicts the superior patient stratification of the Radiomics-T1FSGd grading model observed in this study.

Our models achieved accuracies for the detection of high-grade STS of 0.78 (Radiomics-T1FSGd and Radiomics-T2FS) to 0.83 (Radiomics-combined) with good positive predictive values (PPV) (0.86–0.88) on the expense of negative predictive values (NPV). Thus, classifications of high-grade STS ("positive prediction") provided a more reliable results than the prediction of a low-grade STS. The number of false negative results (i.e. high-grade STS that were classified as low-grade STS) impaired the prognostic validation. The large observed difference between PPV and NPV, as well as the suboptimal patient stratification may be in part explained by the large prevalence of high-grade sarcomas in the validation set (81% of patients). Tumor grading itself achieved significant patient stratification in this skewed validation set only by a close margin ($p = .045$) making a significant patient stratification for surrogate markers, such as the proposed radiomic models, even more difficult. A future external TRIPOD type 4 validation with a more balanced patient data optimally reflecting the true tumor grading distribution may help to assess the impact of the proposed models in terms of classification and prognostic stratification [34].

There are several limitations to this work. In the current study, one could observe a drop in performance for the T1FSGd-based grading models. We applied nested cross validation for an optimally unbiased performance evaluation on the training set while trying to reduce overfitting. Still, reproducibility remains suboptimal with absolute differences in AUC of 0.09 and 0.08 for Radiomics-T1FSGd and Radiomics-combined between training and validation sets, respectively. There are potential explanations for this phenomenon. First, there was a large technical heterogeneity inside and between the two cohorts. In total, 20 MRI scanner types produced by four manufacturers were used across the two cohorts. Thirteen scanner types in the validation set were not present in the training set. The improved testing performance after performing batch correction with an absolute increase in AUC of up to 0.08 may be a sign of the important role of such batch-dependent effects. In contrast, Radiomics-T2FS showed good reproducibility even without batch correction. Secondly, STSs constitute one of the most heterogeneous malignant entities with over 100 histological subtypes [35]. In our study, a direct comparison of histological subtypes revealed a significant difference between training and testing cohort. Given the propensity of radiomics to predict tumor grading and molecular aberrations, the histological subtype may have a substantial influence on the radiomics phenotype. In particular, the amount of contrast enhancement of specific histologies may be an interfering factor, which may also explain that the T2FS-based model without contrast agent showed better reproducibility. In particular, the large proportion of liposarcomas in the training set (39%) may be an explanatory cause. Including histology as biological covariate into ComBat harmonization was not able to improve reproducibility. One solution to these problems may be a future prospective trial which would need to be sufficiently large to achieve a representative distribution of all histological subtypes. If future cohort sizes would achieve a critical number, even better performances may be possible by restricting model building to single histological subtypes.

To conclude, we developed MRI-based radiomic grading models differentiating high-grade from low-grade STS. External validation confirmed classification performance. A radiomic classifier using T2FS sequences appeared to be superior to a T1FSGd-based model predicting tumor grading. Integrating the Radiomics-T2FS model with the AJCC clinical staging system improved prognostic assessment and the clinical net benefit.

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**Declaration of competing interests**

The authors declare no potential conflicts of interest.

**Author contributions**

Jan C. Peeken: Design of the work, collection of data, data analysis, statistical analyses and writing of the manuscript.
Matthew B. Spraker: Collection of data and data analysis.
Carolin Knebel: Collection of data and writing of the manuscript.
Hendrik Dapper: Collection of data.
Daniela Pfeiffer: Data analysis.
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Ahmed Thamer: Collection of data, data analysis.
Mohamed A. Shouman: Collection of data, data analysis.
Armin Ott: Statistical analyses.
Rüdiger von Eisenhart-Rothe: Revision of the manuscript.
Fridtjof Nüsslin: Design of the work and revision of the manuscript.
Nina A. Mayr: Design of the work and revision of the manuscript.
Matthew J. Nyfflot: Design of the work, literature search and revision of the manuscript.
Stephanie E. Combs: Design of the work and revision of the manuscript.

**Appendix A. Supplementary data**

Supplementary data to this article can be found online at https://doi.org/10.1016/j.ebiom.2019.08.059.

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