Texture-Based Analysis of 100 MR Examinations of Head and Neck Tumors – Is It Possible to Discriminate Between Benign and Malignant Masses in a Multicenter Trial?

Zusammenfassung

Ziel: Ziel der Studie war die Auswertung der Texturanalyse in Bezug auf eine mögliche Unterscheidung zwischen benignen und malignen Kopf-Hals-Raumforderungen mittels konventioneller MRT-Sequenzen.


Kernaussagen:
1. Kopf-Hals-Raumforderungen können mittels 2D/3D-Texturanalyse untersucht werden

Abstract

Aim: To evaluate whether texture-based analysis of standard MRI sequences can help in the discrimination between benign and malignant head and neck tumors.

Materials and Methods: The MR images of 100 patients with a histologically clarified head or neck mass, from two different institutions, were analyzed. Texture-based analysis was performed using textural analysis software, with region of interest measurements for 2D and 3D evaluation independently for all axial sequences. COC, RUN, GRA, ARM, and WAV features were calculated for all ROIs. 10 texture feature subsets were used for a linear discriminant analysis, in combination with k-nearest-neighbor classification. Benign and malignant tumors were compared with regard to textural-based values.

Results: There were differences in the images from different field-strength scanners, as well as from different vendors. For the differentiation of benign and malignant tumors, we found differences on STIR and T2-weighted images for 2D, and on contrast-enhanced T1-TSE with fat saturation for 3D evaluation. In a separate analysis of the subgroups 1.5 and 3 Tesla, more discriminating features were found.

Conclusion: Texture-based analysis is a useful tool in the discrimination of benign and malignant tumors when performed on one scanner with the same protocol. We cannot recommend this technique for the use of multicenter studies with clinical data.

Key Points:
1. 2D/3D texture-based analysis can be performed in head and neck tumors
2. Texture-based analysis can differentiate between benign and malignant masses
3. Analyzed MR images should originate from one scanner with an identical protocol

Authors

J. Fruehwald-Pallamar1, J. R. Hesselink2, M. F. Mafee2, L. Holzer-Fruehwald3, C. Czerny1, M. E. Mayerhofer3

Affiliations

1 Department of Biomedical Imaging and Image-guided Therapy, Subdivision of Neuroradiology and Musculoskeletal Radiology, Medical University of Vienna, Austria
2 Department of Radiology, UCSD Medical Center, San Diego, United States
3 Department of Biomedical Imaging and Image-guided Therapy, Medical University of Vienna, Austria

Bibliography

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Correspondence

Frau Dr. Julia Fruehwald-Pallamar
Department of Biomedical Imaging and Image-guided Therapy, Subdivision of Neuroradiology and Musculoskeletal Radiology, Medical University Vienna Waehringer Guertel 18–20
1090 Vienna Austria
Tel.: ++ 43/1/40 40 04 81 80
Fax: ++ 43/1/40 40 04 89 80
julia.fruehwald-pallamar@meduniwien.ac.at

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1. 2D/3D texture-based analysis can be performed in head and neck tumors
2. Texture-based analysis can differentiate between benign and malignant masses
3. Analyzed MR images should originate from one scanner with an identical protocol
2. Es ist möglich benign und maligne Raumforderungen anhand von Texturparametern zu unterscheiden.
3. Die MRT Untersuchung sollte mit gleichem Protokoll auf einem Gerät stattfinden.

**Introduction**

Malignant head and neck tumors account for approximately 3% of all malignancies [1]. Over 85% of head and neck tumors are squamous cell carcinomas (SCC). They are strongly related to tobacco and alcohol abuse, more common in the male population, and mostly located on the mucosal surface. Tumor extent is usually evaluated with cross-sectional imaging.

Texture analysis is a computer-assisted technique that is conducive to the detection and quantification of mathematical patterns called texture features. Those texture features exist in the gray-level distribution of pixels of digital images, which the human eye can only recognize to a limited degree and is unable to quantify. With texture-based analysis, it is possible to distinguish between different types of tissues, and also between healthy and pathologically altered tissues. Texture features derived from the gray-level histogram, the co-occurrence matrix (COC; information about the gray-level value distribution of pairs of pixels, separated by a defined distance, in a given direction), the run-length matrix (RUN; information about runs of pixels with the same gray-level values, in a given direction), the absolute gradient (GRA; information about sudden signal intensity changes in the gray-level values), the auto-regressive model (ARM; description of texture based on the statistical correlation between neighboring pixels), and the wavelet transform (WAV; information about the frequency content of an image within different scales of that image) are used for tissue differentiation [2–5].

Texture analysis of MR images is of particular interest, because this imaging modality offers excellent depiction of anatomic details, high soft-tissue contrast, and allows enhancement of different types of tissues through the use of different pulse sequences, even without the application of contrast media [6, 7].

The aim of the study was to determine whether texture analysis, a manually drawn region-of-interest (ROI) was defined independently for each lesion and MRI sequence, covering the following sequences: 97 T1-TSE, 22 T2-TSE, 46 T2-TSE fs, 31 STIR, T2-TIRM 6000 73 150 320 × 256

<table>
<thead>
<tr>
<th>manufacturer</th>
<th>field strength</th>
<th>sequence (axial)</th>
<th>TR (ms)</th>
<th>TE (ms)</th>
<th>IT (ms)</th>
<th>matrix</th>
</tr>
</thead>
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</tr>
<tr>
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<td>1.5 T</td>
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<tr>
<td></td>
<td>T2-FSE fs</td>
<td>4000 – 6000</td>
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<td>256 × 224</td>
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<td>Ce T1-FSE fs</td>
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<td>320 × 224</td>
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<tr>
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<td>T1-FSE</td>
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<td>7</td>
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<td></td>
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<tr>
<td></td>
<td>T2-FSE fs</td>
<td>4000 – 5000</td>
<td>85</td>
<td>384 × 256</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Materials and Methods**

This retrospective study included patients from two institutions who underwent a routine MR examination, and who subsequently had surgery or biopsy for pathologic correlation. The study was approved by both institutional review boards. A search for patients was performed using a full-text search in the radiology information system between January 2008 and August 2011. The single criterion for inclusion in our study was the presence of one head and neck mass, proven through fine-needle aspiration, biopsy, or operation. Exclusion criteria were the presence of motion artifacts, or a maximal lesion diameter < 5 mm. This 5 mm cut-off value was chosen to minimize the influence of partial volume effects, which might distort the true tissue-specific image texture. Based on the above-defined criteria, 100 patients (50 per institution; 55 male, 45 female) with a mean age of 50.9 years (ranging from 12 to 83 years) were included.

The MR examinations were performed on different MR scanners; 1.5 T GE (General Electric) and Siemens scanners and a 3 T GE scanner at site A, and a 1.5 T and 3 T Siemens scanner and a 3 T Philips scanner at site B. The standard examination protocols also varied somewhat. At site A, the standard protocol was: axial T1 TSE (turbo spin echo); T2 TSE fs (fat saturation) and coronal T1; and axial and coronal T1 TSE with fat saturation after gadolinium administration. The standard protocol of site B was: axial and coronal STIR (short tau inversion recovery) or T2-TIRM; axial T1-TSE before and after contrast administration; and coronal T1-TSE fs after contrast administration (sequence parameters are listed in **Table 1**). We decided to evaluate only the axial sequences (T1-TSE, STIR, T2-TSE fs, contrast-enhanced T1-TSE, and T1-TSE fs). Due to the retrospective character of the study, some sequences were not available in some patients. In total, we evaluated the following sequences: 97 T1-TSE, 22 T2-TSE, 46 T2-TSE fs, 31 STIR, 48 contrast-enhanced (ce) T1-TSE, and 43 with fat saturation.

Texture analysis was performed independently for all axial sequences (STIR, T2-TSE fs, T1-TSE before and after i.v. contrast administration (with fat saturation)). First, for the 2 D texture analysis, a manually drawn region-of-interest (ROI) was defined independently for each lesion and MRI sequence, covering the

**Table 1** Sequence parameters.

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entire lesion on the image that depicted the lesion at its greatest diameter. A semi-automatic, active computer algorithm using the default setting and automatic fill-in defined the ROI. Then, for the 3D texture analysis, the tumor was encircled on every slice visible on every axial sequence in the same manner. Gray-level histogram features, co-occurrence matrix (COC) and run-length matrix (RUN) features were calculated for all 2D and 3D ROIs, using the popular texture analysis software MaZda (version 4.7, available at http://www.eletel.pl/odz.pl/mazda/) [8]. Additionally for all 2D ROIs absolute gradient (GRA), autoregressive model (ARM), and wavelet transform (WAV) features were calculated by the same program (Table 2).

First, we wanted to investigate whether field strength influenced the performance of texture analysis on 1.5 T versus 3 T. In order to analyze this influence, we chose to evaluate and compare the texture features within our largest homogeneous group, the squamous cell carcinomas. With this group, we also analyzed whether there were differences when performing texture analysis on the MR images from different vendors (GE, Siemens, Philips).

We then tested our primary hypothesis regarding whether we could find texture features that would distinguish between benign and malignant head and neck tumors. To identify the most valuable texture features for distinguishing between benign and malignant lesions, three subsets of 10 texture features were extracted, independently for T1-weighted, T2-TSE (±fs), STIR, and contrast-enhanced T1-weighted images (±fs), based on Fisher coefficients (ratio of between-class to within-class variance), minimization of both classification error probability and average correlation coefficients (POE+ACC), and mutual information (MI) coefficients, which measure the dependence between two or more random variable coefficients [2, 9]. This extraction was performed with the program, B11, which is integrated in the MaZda software.

For computer-assisted differentiation between benign and malignant head and neck tumors, based on the previously calculated texture feature subsets (Fisher, POE+ACC, and MI), we used linear discriminant analysis (LDA) in combination with k-nearest-neighbor classification (k-NN). K-NN calculates the rate of misclassified data vectors by comparison with the previously defined true class affiliations. The k-NN classifier implemented in the MaZda software uses the leave-one-out testing technique for this task. Thus, no separation of datasets into a training and test dataset is required. The rates of correctly classified data vectors were used as the primary outcome variables. This strategy of texture-based lesion classification has been used in previously published papers [3–5, 10].

**Results**

Of the 100 patients in our study with a head and neck mass, 54 patients (33 male, 21 female) were diagnosed with a malignant mass and 46 patients with a benign mass (24 female, 22 male). The largest histologically homogeneous group was the patients

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**Table 2** List of texture features and abbreviations (as used in Table 4, 5).  

<table>
<thead>
<tr>
<th>Gray-level histogram</th>
<th>mean, variance, skewness, kurtosis (percentiles: 1, 10, 50, 90, 99 %)</th>
<th>mean (3 D) variance (3 D) skewness (3 D) kurtosis (3 D) perc.01 %, Perc.10 %, Perc.50 %, Perc.90 %, Perc.99 % (3 D)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Co-occurrence matrix</td>
<td>S(2.2)AngScMom, S(3.3)AngScMom, S(1.1)AngScMom, S(5.5)AngScMom, S(0.2)AngScMom, S(3.0)Contrast, S(1.3)Contrast, S(4.4)Contrast, S(5.0)Contrast, S(1.0)Contrast, S(5.0)Correlat, S(5.5)Correlat, S(2.2)Correlat, S(1.1)Correlat, S(5.0)Correlat, S(1.3)Correlat, S(3.0)Entropy, S(5.5)Entropy, S(2.2)Entropy, S(4.4)Entropy, S(5.0)Entropy, S(1.3)Entropy, S(1.0)SumEntrop, S(0.2)SumEntrop, S(4.0)SumEntrop, S(3.0)SumEntrop, S(3.0)SumOfSqss, S(4.0)SumOfSqss, S(5.0)SumOfSqss, S(5.5)SumOfSqss, S(2.2)SumAverg, S(3.3)SumAverg, S(0.3)SumAverg, S(0.5)SumAverg, S(0.4)SumAverg, S(1.1)SumVarncc, S(3.0)SumVarncc, S(5.5)SumVarncc, S(0.3)INVdMom, S(1.0)INVdMom, S(4.4)INVdMom, S(5.0)INVdMom, S(0.1)INVdMom, S(5.5)DIFEntrp, S(4.4)DIFEntrp, S(1.1)DIFEntrp, S(0.1)DIFEntrp, S(2.0)DIFEntrp, S(3.3)DIFVarncc, S(1.1)DIFVarncc, S(2.2)DIFVarncc, S(5.5)DIFVarncc, S(2.0)DIFVarncc</td>
<td></td>
</tr>
<tr>
<td>Absolute gradient (GRA)</td>
<td>GrMean, GrVariance, GrSkewness, GrKurtosis, GrNonZeros</td>
<td></td>
</tr>
<tr>
<td>Autoregressive model (ARM)</td>
<td>theta 1 to 4, sigma, Teta1, Teta2, Teta3, Teta4</td>
<td></td>
</tr>
<tr>
<td>Wavelet transform (WAV)</td>
<td>energies of wavelet transform coefficients in sub-bands LLH, HLH, HH (for three subsampling factors)</td>
<td>WavEnLL_s-1, WavEnLL_s-2, WavEnLL_s-3, WavEnLL_s-4, WavEnLH_s-1, WavEnLH_s-2, WavEnLH_s-3, WavEnHL_s-1, WavEnHL_s-2, WavEnHL_s-3, WavEmHH_s-1, WavEmHH_s-2, WavEmHH_s-3</td>
</tr>
</tbody>
</table>

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...
diagnosed with (SCC) squamous cell carcinoma (n = 31; 21 male, 10 female; Table 3).

**Table 3** Pathologic subgroups.

<table>
<thead>
<tr>
<th>histologic subgroup</th>
<th>number of patients</th>
</tr>
</thead>
<tbody>
<tr>
<td>benign tumors n = 46</td>
<td></td>
</tr>
<tr>
<td>cysts</td>
<td>8</td>
</tr>
<tr>
<td>inflammatory mass/abscess</td>
<td>5</td>
</tr>
<tr>
<td>parotid tumor</td>
<td>9</td>
</tr>
<tr>
<td>glomus tumor</td>
<td>9</td>
</tr>
<tr>
<td>vascular lesion/malformation</td>
<td>5</td>
</tr>
<tr>
<td>schwannoma</td>
<td>4</td>
</tr>
<tr>
<td>other</td>
<td>6</td>
</tr>
<tr>
<td>malignant tumors n = 54</td>
<td></td>
</tr>
<tr>
<td>squamous cell carcinoma</td>
<td>31</td>
</tr>
<tr>
<td>lymphoma</td>
<td>8</td>
</tr>
<tr>
<td>adenoid cystic carcinoma</td>
<td>5</td>
</tr>
<tr>
<td>adenocarcinoma</td>
<td>4</td>
</tr>
<tr>
<td>other</td>
<td>6</td>
</tr>
</tbody>
</table>

**SCC – comparison of vendors and field strength**

Considerable differences in terms of texture features were observed between the squamous cell carcinomas at 1.5 and 3 Tesla. Discrimination based on 2D texture features extracted from T1-weighted sequences produced misclassification numbers of 12.9 % for POE+ACC, and 19.35 % for MI. We found a low misclassification rate only for MI (19.35 %). Similar results were found in the analysis of the T1-weighted sequences with the 3D ROI (correctly classified data vectors [percentages] and the list of the 10 extracted features are listed in Table 4). For the discrimination of SCC, mutual information coefficients proved to be the most effective method.

**Table 4** List of correctly classified data vectors and the texture feature subset best suited for the discrimination of SCC at 1.5 and 3.0 Tesla.

<table>
<thead>
<tr>
<th>sequence</th>
<th>fisher</th>
<th>POE+ACC</th>
<th>MI</th>
</tr>
</thead>
<tbody>
<tr>
<td>2 D T1-TSE</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>features</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 Perc.01 %</td>
<td>1 Vertl_RLNonUni</td>
<td>1 S(4.4)DifEntrp</td>
<td>27/31 or 87.1 %</td>
</tr>
<tr>
<td>2 S(5.5)DifEntrp</td>
<td>2 Vertl_RLNonUni</td>
<td>2 S(1,-1)DifEntrp</td>
<td>25/31 or 80.65 %</td>
</tr>
<tr>
<td>3 Perc.10 %</td>
<td>3 Vertl_ShrtRemp</td>
<td>3 S(3.0)Entropy</td>
<td>25/31 or 80.65 %</td>
</tr>
<tr>
<td>4 Mean</td>
<td>4 Mean</td>
<td>4 S(0.3)SumAverg</td>
<td>25/31 or 80.65 %</td>
</tr>
<tr>
<td>5 Perc.90 %</td>
<td>5 Vertl_ShrtRemp</td>
<td>5 S(0.5)SumAverg</td>
<td>25/31 or 80.65 %</td>
</tr>
<tr>
<td>6 Perc.50 %</td>
<td>6 Vertl_ShrtRemp</td>
<td>6 S(0.3)InvMom</td>
<td>25/31 or 80.65 %</td>
</tr>
<tr>
<td>7 Perc.99 %</td>
<td>7 Vertl_ShrtRemp</td>
<td>7 135dr_Fraction</td>
<td>25/31 or 80.65 %</td>
</tr>
<tr>
<td>8 Vertl_RLNonUni</td>
<td>8 S(4.4)DifEntrp</td>
<td>8 135dr_ShrtRemp</td>
<td>25/31 or 80.65 %</td>
</tr>
<tr>
<td>9 S(4.4)DifEntrp</td>
<td>9 Vertl_RLNonUni</td>
<td>9 135dr_ShrtRemp</td>
<td>25/31 or 80.65 %</td>
</tr>
<tr>
<td>10 Vertl_ShrtRemp</td>
<td>10 Vertl_RLNonUni</td>
<td>10 135dr_ShrtRemp</td>
<td>25/31 or 80.65 %</td>
</tr>
<tr>
<td>without gray-level histogram features</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>24/31 or 77.42 %</td>
<td>21/31 or 67.7 %</td>
<td>25/31 or 80.65 %</td>
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</tr>
<tr>
<td>3 D T1-TSE</td>
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</tr>
<tr>
<td>features</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 S(5.5)DifEntrp</td>
<td>1 Vertl_GLevNonU</td>
<td>1 S(4.4)DifEntrp</td>
<td>24/31 or 77.78 %</td>
</tr>
<tr>
<td>2 Vertl_RLNonUni</td>
<td>2 Vertl_GLevNonU</td>
<td>2 S(2,-2)AngScMom</td>
<td>24/31 or 77.78 %</td>
</tr>
<tr>
<td>3 S(4.4)DifEntrp</td>
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<td>3 S(3.0)Entropy</td>
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<td>6 S(5.0)Correlat</td>
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<td>7 S(3.0)Entropy</td>
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<td>8 45dgr_ShrtRemp</td>
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<td>24/31 or 77.78 %</td>
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<tr>
<td>9 S(3.3)DifVarnc</td>
<td>9 Vertl_GLevNonU</td>
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<td>24/31 or 77.78 %</td>
</tr>
<tr>
<td>10 Vertl_Fraction</td>
<td>10 Vertl_Fraction</td>
<td>10 135dr_ShrtRemp</td>
<td>24/31 or 77.78 %</td>
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</table>

<table>
<thead>
<tr>
<th>T2-TSE fs</th>
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<td>T1-TSE</td>
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<td></td>
</tr>
<tr>
<td>features</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 S(5.5)DifEntrp</td>
<td>1 Vertl_GLevNonU</td>
<td>1 S(4.4)DifEntrp</td>
<td>14/18 or 77.78 %</td>
</tr>
<tr>
<td>2 45dgr_Fraction</td>
<td>2 S(2,-2)AngScMom</td>
<td>2 S(1,-1)DifEntrp</td>
<td>14/18 or 77.78 %</td>
</tr>
<tr>
<td>3 S(5,-5)Correlat</td>
<td>3 Vertl_ShrtRemp</td>
<td>3 S(3.0)Entropy</td>
<td>14/18 or 77.78 %</td>
</tr>
<tr>
<td>4 S(2,-2)Correlat</td>
<td>4 Variance</td>
<td>4 S(1.0)DifEntrp</td>
<td>14/18 or 77.78 %</td>
</tr>
<tr>
<td>5 S(4.5)Entropy</td>
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<td>14/18 or 77.78 %</td>
</tr>
<tr>
<td>6 Vertl_LngRemph</td>
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<tr>
<td>7 S(5,-5)DifEntrp</td>
<td>7 Vertl_GLevNonU</td>
<td>7 S(1.0)Entropy</td>
<td>14/18 or 77.78 %</td>
</tr>
<tr>
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<td>8 Vertl_GLevNonU</td>
<td>8 S(1.0)DifEntrp</td>
<td>14/18 or 77.78 %</td>
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<td>14/18 or 77.78 %</td>
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<td>1 S(4.4)DifEntrp</td>
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<td>2 Vertl_LngRemph</td>
<td>2 Vertl_Fraction</td>
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<td>3 S(3.0)Entropy</td>
<td>3 Vertl_LngRemph</td>
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<td>4 Vertl_Fraction</td>
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<tr>
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<td>6 Vertl_LngRemph</td>
<td>6 S(2,-2)Entropy</td>
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<td>10 Vertl_Fraction</td>
<td>10 S(0.1)DifEntrp</td>
<td>24/31 or 77.78 %</td>
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We also found textural differences between the three different vendors (GE, Siemens, and Philips) for the 2D ROIs on the T1-weighted sequence, with the following correctly classified data vectors: 26/31 or 83.87% for Fisher (41.94% without gray-level histogram features); 24/31 or 77.42% for POE+ACC, and 28/31 or 90.32% for MI (80.65% without gray-level histogram features).

**Discrimination of benign and malignant tumors**

For the discrimination of benign and malignant lesions, STIR and T2-weighted images contained the most relevant texture features for 2D evaluation. For 3D texture-based analysis, only contrast-enhanced, T1-weighted images with fat saturation had a low misclassification rate. ○ Table 5 shows the results for the LDA of the comparison between benign and malignant tumors.

### Table 5  List of correctly classified data vectors and the texture feature subset best suited for the discrimination of benign and malignant tumors.

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<th>MI</th>
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<td>3 Variance</td>
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<td>9 S(2.2)DifVarnc</td>
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<td>10 S(3.3)DifVarnc</td>
<td>10 S(4,-4)InvDfMom</td>
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<td>STIR</td>
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<td>20/31 or 64.52%</td>
<td>25/31 or 80.65%</td>
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<td>3 Variance</td>
<td>3 WavEnLH_s-1</td>
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<td>10 S(3.3)Correlat</td>
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<td>36/48 or 75%</td>
<td>27/48 or 56.25%</td>
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<td>22/22 or 100%</td>
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We then decided to analyze the 2 D ROI of benign and malignant tumors at 1.5 and 3.0 Tesla separately. This analysis showed better results in the discrimination between benign and malignant tumors. For the T1-weighted images, we found 41/49 or 83.67 % correctly classified data vectors at 3 T with MI. The contrast-enhanced, T1-weighted images with fat saturation at 1.5 T had 86.67 % (or 26/30) correctly classified data vectors. For the T2-weighted images with fat saturation (fs), all vectors were correctly classified at 3 T, with 81.48 % (Fisher), and, at 1.5 T, 92.59 % (POE + ACC and MI) were correctly classified (n = 27).

Additionally, we analyzed the 2 D ROI on axial STIR images of a single 3 T unit (n = 28). All vectors were correctly classified in 71.43 % vs. 68.82 % (n = 31) for Fisher, POE+ACC and MI – in detail: 16/28 for Fisher; 18/28 for POE+ACC, and 22/28 for MI.

Discussion

In contrast to the published literature about the application of texture-based analysis in phantoms and in patients examined on one scanner with the same protocol, our data show that, when using different MR protocols on different MR scanners (field-strength, vendor), texture-based analysis is not practical for the differentiation between benign and malignant head and neck tumors. Although we found discriminating features for STIR and T2-weighted images, the number of correctly classified data vectors was not sufficient to implement this technique in the routine evaluation of MR images.

Several studies have shown the successful use of texture-based analysis in the brain [11, 12], the liver [3, 10], and muscle [13]. Herlidu-Meme et al. performed MR imaging at three sites, but with the same brand and field-strength MR unit [14]. Jirak et al. performed phantom measurements on six different MR scanners, all using identical MR parameters for the acquisition protocol [15]. In a recently published study, Fruehwald-Pallamar et al. were able to discriminate between benign and malignant parotid tumors based on texture analysis [5] – notably, using images obtained using a single MR unit and identical protocol.

Contrary to the above described successful applications of texture analysis with (more or less) uniform scanning conditions, previous studies indicated considerable influence of protocol heterogeneity. In two articles Lerski et al. reported that texture-based information extracted from foam test objects was susceptible to the choice of MR protocol and scanner [16, 17]. Mayerhoefer et al. showed in a phantom study that texture features (especially COC and RUN) are sensitive to variations in the scanned protocol (matrix size, TR, TE, and number of acquisitions) [7].

To our knowledge, texture-based analysis has not been applied for the analysis of different tissue types (different kinds of head and neck masses) in a multicenter study. In the present study, routine MR images were obtained on several MR units, with similar imaging protocols (plane, and the use of fat saturation), but with each center using its own acquisition parameters. This was done to determine the clinical value of this technique and to analyze which texture feature subset and extraction method would perform best under given conditions. Therefore, we also tried to acquire a large collective of 100 patients.

Our clinical data resemble the phantom data from the above-mentioned study by Lerski et al. because the texture features extracted from SCC clearly differed between examinations performed at different field strengths, and also between images obtained from scanners by different vendors [16]. For instance, our classifier was able to differentiate between the SCC at 1.5 and the SCC at 3 Tesla in almost 90 % of the cases, based on texture features extracted from T2 TSE fs images. It is quite possible that the strong influence of acquisition parameters affected the texture-based discrimination of benign and malignant head and neck tumors in our study. When performing the analysis with the data of one scanner (benign vs. malignant lesions on one 3 T unit), we were able to increase the correctly classified data vectors by 3 % for axial STIR images. Our results were generally less favorable than those reported by Fruehwald-Pallamar et al. [5] in a single-center study using 2 D texture features. Notably, our results indicate that 3 D texture features are not superior to a 2 D evaluation of a single slice for head and neck masses—this topic has not been evaluated thus far in a clinical setting. For a possible future introduction into routine clinical practice, and also for clinical trials involving texture analysis, we recommend that only data from a single scanner are used, in order to avoid an incalculable influence from hardware specifications. Furthermore, 2 D evaluation is less time-consuming, and therefore, more practical, while providing results comparable to 3 D evaluation, where every slice of the tumor has to be processed.

We note the limitations of our study: There were minor acquisition parameter differences, even between comparable pulse sequences obtained by the different scanners, as mentioned above.

In conclusion, texture-based analysis has the potential to help with the discrimination of benign and malignant head and neck tumors when performed on the same scanner with an identical protocol. We cannot recommend the application of texture-based analysis in a multicenter study with different types of scanners and varying acquisition parameters.

Clinical relevance

Since morphological sequences alone often do not allow a correct diagnosis of different types of tumors in the head-neck region, it is of great help for radiologists to have additional information to generate the correct diagnosis. Texture-based analysis is a tool, which has the potential to help with the discrimination between benign and malignant head and neck tumors, provided that the examinations take place on one specific machine with a defined protocol.

Acknowledgments

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