# Automatic Segmentation of Necrosis Zones after Radiofrequency Ablation of Spinal Metastases

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Abstract: In this work, we propose an automatic deep learning-based approach to segment necrotizing tissue (necrosis zones) after radiofrequency ablations (RFA) of spinal metastases in follow-up Magnetic Resonance (MR) images. While the manual segmentation of those necrosis zones is challenging and time consuming, it is a crucial step to assess, whether a preceding therapy using RFA was successful and to what extent, i.e., to quantitatively evaluate how much of the metastasis was necrotized throughout the therapy. Therefore, we trained a U-Net like deep neural network on 26 clinical cases (and various augmentations of those), where each case had an associated contrast enhanced  $T_1$ -weighted as well as a  $T_2$ -weighted MR sequence. We evaluated the proposed approach on both sequences separately as well as in a combined setting and report Dice coefficients, sensitivity-, and specificity rates for the automatic segmentations. A Dice coefficient of up to 77.2 % indicates promising segmentation quality, if compared to related work and similar segmentation tasks. To the best of our knowledge, this is the first work to tackle the problem of automatic segmentation of necrosis zones in MR images and therefore lacks comparability with related works. However, our best results are somewhat superior to semi-automatic approaches of liver metastases segmentation, which might be considered a problem of similar complexity.

# **1 INTRODUCTION**

The life expectancy has increased steadily over the last decades, resulting in a lifetime gain, which, in addition to all its benefits, also holds risks. Age-related diseases like cardiovascular diseases, as well as cancer and cancer induced malicious metastases occur more and more frequently. Due to improved diagnostic procedures and medical treatments, the survival time of most malicious carcinomata has increased. This medical success story unfortunately again has a drawback, as the probability to develop metastases raises. Bone metastases are the third most likely and a vast majority are located in the spine (Harrington, 1986; Wong et al., 1990). These spinal metastases can evoke vigorous pain by fractures, bruises, spinal cord and nerve root compressions or neurologic deficits and therefore, detrimentally affect the quality of life (Klimo and Schmidt, 2004). Besides, externalbeam radiation the method of choice are ablative approaches like radiofrequency ablation (RFA) (Rosenthal et al., 1998; Dupuy et al., 2010). The underlying princible is based on tissue heating and necrotization by molecular firction due to high frequency current phase changes at the tip of a minimally invasive inserted applicator. Heat dissipation leads to a growing necrotized zone around the needle tip, with the purpose of covering as much of the metastasis as possible. MR images are taken for both diagnostic purposes and post-interventional follow-ups, due to their high soft tissue contrast and spatial resolution as well as the possibility to utilize contrast agents. To assess the treatment outcome, i.e. the completeness of metastasis ablation it is essential to evaluate the congruence of both the ablation zone and the former metastasis via distance and volume overlap measures. Therefore, the segmentation of the necrosis zone is among others a pivotal step towards an automatised, quantitative and objective treatment outcome validation of RFAs of spinal metastases. Besides, such segmentations could yield additional insights for a better understanding in tumor reoccurrence and to further improve ablation protocols and procedures. Currently, it is not part of the clinical routine, due to the very time-consuming manually contouring done slice-wise, which makes it hardly applicable or suitable. Furthermore, some specific aspects of this issue make this a highly challenging and ambitious task,



Figure 1: Original  $T_1$ -weighted pre- (upper row) and contrast-enhanced  $T_1$ -weighted post-interventional (lower row) MR images of exemplary patient cases with spinal metastases and corresponding necrosis zones after RF ablations (arrows).

e.g. hardly noticeable intensity differences between necrosis and remaining metastasis, inflammation processes caused by the coagulation, possible hematoma as well as a high shape variability. The latter depends on the former metastasis shape, tissue specific heat dissipation, and adjacent anatomical structures (see Fig. 1).

Currently, there are only few works regarding RF ablation zone segmentation. To our best knowledge, all published research papers focused on liver tumour treatment in Computed Tomography (CT) imaging, none of them dealt with spine RFA necrosis zones or MR imaging. Therefore, the following state-ofthe-art analysis cannot be or only partially transferred to ablation zone segmentation in spine MR imaging. Passera et al. (Passera et al., 2013) proposed a live-wire algorithm and Fuzzy C-Mean clustering to segment ablation zones in post-interventional RFA CT scans of the liver. The semi-automatic and userguided approach was applied as a 2D slice-wise segmentation and took on average approx. 10 min for a necrosis zone. Similar to the work by Passera et al., McCreedy et al. (McCreedy et al., 2006) likewise presented a 2D live-wire based approach for CT imaging, embedded in a RFA registration, segmentation, and fusion tool. They described the segmentation step merely superficial and did not present any quantitative results. Weihusen et al. (Weihusen et al., 2010) proposed a workflow oriented software support for CT image guided RFA of focal liver malignancies. As a treatment outcome validation tool, they also included a semi-automatic necrosis zone segmentation, based on a morphological region growing algorithm. Again, no quantitative results were stated.

Another computer-aided analytic tool was developed by Bricault et al. (Bricault et al., 2006), which focused on assessing local recurrences of liver metastases after RFA treatment in CT. For this purpose, a semi-automatic 3D segmentation approach based on a watershed algorithm was implemented. On average, the segmentation took approx. 4 min, but accuracy results were not stated. A semi-automatic graph-cuts based approach was proposed by Egger et al. (Egger et al., 2015) to segment liver tumors in CT imaging. They achieved accuracies of 77 % Dice on their 12 patient comprising dataset and had computational times of only a few seconds. In conclusion, it remains unclear, why most of the above mentioned works did not state any segmentation accuracy results whatsoever, since the segmentation of the ablation zone is a cru-



Figure 2: The U-Net structure used for multi-modal 2D image input, with convolutional layers including batch normalisation, strided convolutions for downsampling and upsampling layers. The architecture for three-dimensional input is analogous to the one shown above. A significant difference between the two variants is the number of trainable parameters, which is about 2.85 times higher in the 3D case.

cial step towards assessing the efficacy of RFA treatment of metastases and tumours. Furthermore, the previous studies focused only on semi-automatic approaches for post-interventional CT imaging of liver tumor ablations. The main objective of this study was to implement an automatic, deep-learning based segmentation approach for follow-up MR scans of RFAs of spinal metastases and to assess, which image input w.r.t. dimension and MR sequence for a Convolutional Neural Network (CNN) is most suitable.

## 2 MATERIALS AND METHOD

### 2.1 Image Data

We assembled a dataset of 26 follow-up MR images of patients who underwent radiofrequency ablations of spinal metastases. These images were acquired between 1 and 3 days after the intervention including, among others, sagittal contrast-enhanced  $T_1$ -weighted and native  $T_2$ -weighted MRI sequences. Both sequences were chosen, since they are most commonly used for visual treatment outcome validation by neuroradiologists in this particular case. Due to multiple MR scanners, the specific acquisition parameters, e.g. magnetic field strength or repetition time, varied within our dataset. The scan resolution within a slice ranged between the individual cases from 0.45 mm to 1.25 mm, the spacing between adjacent slices from 3.3 mm to 4.8 mm. The image volumes of each patient case were pre-processed by a cubic interpolation between the original number of slices (between 15 and 25) to a fixed number of 64 to yield almost isotropic spatial resolution and simplify any following processing steps. An experienced neuroradiologists manually contoured each necrosis slice-wise, thus, the input data could be applied as individual slices or as

patient-wise volumes to our networks.

#### 2.2 Augmentation

Due to our relatively small amount of available data, we extensively augmented each of the 26 original MRI volumes using the following techniques:

- *Gaussian blur:* The images were burred with a Gaussian filter with σ in the range from 0 to 0.5.
- Gamma transformation: Gamma transformations with γ in the range from 0.5 to 2 were applied to modify image intensities.
- *Mirroring:* Each patient volume was flipped in all directions. We included vertical flips, i.e. cranio-caudal, even though it may appear inappropriate, it had proven to be advantageous for the final results due to the avoidance of fast overfitting.
- *Scaling:* The image volumes were scaled with randomly chosen factors between 0.6 and 1.4.
- *Rotation:* Rotations were applied to the image volumes in the range of  $\pm 30^{\circ}$  around the transversal axis and between  $\pm 20^{\circ}$  around the sagittal axis.
- *Elastic deformations:* Random displacement fields with subsequently Gaussian smoothing the grid with a  $\sigma$  ranging between 0 and 0.3 were used to elastically deform the image volumes (cf. (Ronneberger et al., 2015)).
- *Translation:* Finally, each patient volume was translated in a random cropping manner within a range of  $\pm 20$  *voxels* in sagittal and vertical direction w.r.t. the center of the necrosis  $m_c$  and subsequently cropped to patches of the fixed size of  $128 \times 128 \times 64$  voxels.

After the augmentation each image volume patch was whitened by mean subtraction and a subsequent

division by the standard deviation. It was ensured that each patch contained at least fractions of necrotizised tissue. Since we used stratified 6-fold crossvalidation with 26 patient cases, we used a 21/5 (training/validation) split for 2 folds and 22/4 for the remaining four. Each patient volume within the training set was augmented 1800 times, yielding in total 37,800 volumetric respectively 2,419,200 crosssection training samples for both 21/5-split folds and 39,600 respectively 2,534,400 for the remaining folds.

### 2.3 CNN architecture

We used minimally modified U-Nets implemented in Keras and Tensorflow, since the commonly used U-Net architecture proposed by Ronneberger et al. (Ronneberger et al., 2015) still represents the state-ofthe-art regarding various medical segmentation tasks (Isensee et al., 2019) (see Fig. 2). The individual networks differ in the processing of either 2D or 3D data and the incorporation of either individual MR sequences or multimodal input. Therefore, we applied 2D patches of size  $128 \times 128$  pixels or volumes of size  $128 \times 128 \times 64$  voxels to our networks with either one modality channel or two. The convolutional layers had a kernel size of  $3 \times 3$  ( $\times 3$ ) except the last one, which applies a  $1 \times 1$  (×1) kernel to reduce the dimensionality to the desired output size. A batch normalization followed each convolutional layer. Strided convolutions (stride of 2) were applied for downsampling the image patches. Simplified upsampling layers replaced the commonly used up-convolutions, since it have been found to be equally effective, while being less computationally expensive (Isensee et al., 2017). The activation function of every convolutional layer was a Rectified Linear Unit (ReLU), except the last one again, where a sigmoid function resulted in values between 0 and 1 were. The above mentioned multimodal image input was incorporated in the most straightforward way, i.e. each MRI sequence was represented by an input channel. We decided to use a single epoch, while the number of iterations was equal to the number of available samples, since we did not alter any training specific parameters. Furthermore, we used Tversky Loss (TL) as proposed by Salehi et al. (Salehi et al., 2017) as a loss function, which represents a modified form of the Tversky index (Tversky, 1977) and is defined as

$$TL(\alpha,\beta) = \frac{2\sum_{i=1}^{N} (r_{0i}p_{0i})}{\sum_{i=1}^{N} (r_{0i}p_{0i}) + \alpha \sum_{i=1}^{N} (r_{1i}p_{0i}) + \beta \sum_{i=1}^{N} (r_{0i}p_{1i})}$$
(1)

where  $p_{0i}$  is the probability for a voxel i to be labeled as necrosis zone and  $p_{1i}$  as background. For a necrosis zone voxel  $r_{0i}$  is 1 and for a background voxel  $r_{0i}$  is 0, vice verse for  $r_{1i}$ .  $\alpha$  and  $\beta$  balance the penalities for false positives and false negatives. Adam (Kingma and Ba, 2014) was applied as an optimizer with a learning rate of 0.001. Mini-batch size was 2 samples for volumetric and 32 for slice-wise input data. Finally, to produce binary output images a threshold of 0.5 was applied.

#### 2.4 Experimental Design

We performed multiple experiments with variing network configurations, i.e. individual or multimodal image input as well as 2D and 3D images. The basic network architecture remained unchanged, in order to largely exclude further influencing factors, e.g. by varying layer numbers or kernel sizes. Our training scheme consisted of stratified 6-fold cross-validation over disjunct subsets of either five or four patients per validation set. The results stated in the following represent the average of all 6 cross-validation folds. Since we did not base any training and design decisions on intermediate validation results (no lookahead bias) and due to our limited dataset, we have decided against a separate test set, as it would result in too few samples for a promising training.

#### 2.5 Evaluation

Expertly annotated necrosis zone segmentations were produced using co-registered MR sequences of each patient within a synchronized viewer. The network described in Section 2.3 was applied to the image data. To quantify our results, we used Dice similarity coefficients to measure the percentage of volume overlap, as well as voxel-wise sensitivity (true positive rate, TPR) and specificity (true negative rate, TNR), since some of the related work used both as quality measurements. The above mentioned are defined as follows:

$$Dice = \frac{2 |R_1 \cap P_1|}{|R_1| + |P_1|}, \ TPR = \frac{|R_1 \cap P_1|}{|R_1|}, \ TNR = \frac{|R_0 \cap P_0|}{|R_0|}$$

with  $R_1$  and  $P_1$  as foreground voxels of reference and prediction and analogously,  $R_0$  and  $P_0$  as the cor-

		2D				3D		
		$ceT_1$	$T_2$	$[ceT_1+T_2]$	$ceT_1$	$T_2$	$[ceT_1+T_2]$	
Dice [%]	mean	76.7	62.2	77.2	72.7	60.4	75.9	
	median	83.5	65.9	82.3	78.1	63.3	80.7	
	std	19.0	21.7	15.6	18.7	27.4	17.2	
	maan	<b>Q1</b> /	60.2	81.6	77 7	63.1	77 8	
Sensitivity [%]	median	84.5	76.0	81.0	86.2	71 1	82.6	
	std	17.1	20.3	15.4	21.5	26.5	19.6	
Specificity [%]	mean	99.2	98.8	99.2	99.1	98.7	99.2	
	median	99.6	99.1	99.5	99.4	99.0	99.6	
	std	0.9	1.0	0.8	0.9	1.3	0.9	

Table 1: Experimental results for each input configuration depending on the used modalities (*contrast – enhanced T*<sub>1</sub>-,  $T_2$ -weighted MRI sequences), as well as a slice-wise (2D) or volume (3D) processing.

responding background voxels. However, sensitivity and specificity are not commonly used to evaluate medical image segmentations, since they are highly sensitive to a segment's size (Taha and Hanbury, 2015). The given results were generated exclusively on patient volumes, even if the segmentations with 2D input were predicted slice-wise. Thus, the 2D predictions were merged patient-wise.

### **3** RESULTS AND DISCUSSION

Table 1 shows the results of our experiments. We achieved Dice scores up to  $77.2 \pm 15.6$  % and sensitivity rates up to  $86.4 \pm 15.4$  %, depending on the applied MR sequences and input dimension. In this study a segmentation approach for necrosis zones in spine MR images was presented. With regards to clinical applicability, the accuracy, automatization, and computational time are of primary importance. Manual segmentation procedures are time-consuming and tedious due to their limitation to slice-by-slice processing and require up to 10 min per necrosis zone. That is one of the reasons, such segmentations are not currently integrated into the clinical routine. Therefore, an automatic and fast apporach can overcome the limitations of manual procedures and play a decisive role in improving treatment outcome validation and ablation protocols. Our proposed method fulfills these requirements. Since this is (to our best knowledge) the first study to tackle necrosis zone segmentation in spine MR images, it is difficult to compare our results with related work. Furthermore, almost all relevant works regarding necrosis zone segmentation did not state any quantitative results, except the work of Egger et al. (Egger et al., 2015). Our best results were on par with the results of their semiautomatic approach (77.2 % vs. 77.0 %), but our method did not require any user-interaction. Due to the lack of directly related work, it might be interesting to take works of automatic lesion segmentation as a similar issue into consideration. Chmelik et al. (Chmelik et al., 2018) adapted a CNN to vertebral metastases segmentation in CT images. They stated a voxel-wise sensitivity rate of 74 % for sclerotic and 71 % for lytic lesions as well as a specificity rate of 88 % (sclerotic) and 82 % (lytic). Hille et al. (Hille et al., 2019) applied a CNN to segment spinal metastases in  $T_1$ - and  $T_2$ -weighted diagnostic MR images and achieved Dice scores up to 73.8 %. Although, our results were superior to those of the mentioned works, it is worth mentioning, that comparability is only possible to a very limited extent. Besides the different imaging method used by Chmelik et al., the segmentation of spinal metastases represents a rather more ambitious task due to the high appearance variability depending on metastatic origin and type (lytic, sclerotic or mixed). Nonetheless, the segmentation of necrosis zones in spine MR imaging is hampered by similar difficulties, as there is a variety of anatomical structures with high image contrasts, similar intensities, and textures in close proximity. Furthermore, the necrosis zones are in some cases difficult to distinguish from remaining metastases and inflammation processes or possible hematoma could overlap with the necrosis zone (see Fig. 4, lower row).

With regards to the applied MR sequence, we achieved the best results with either the contrastenhanced  $T_1$ -weighted data alone or if it was part of multimodal image input (see Fig. 3). This could most likely be attributed to the predominantly high image contrasts between necrosis and surrounding



Figure 3: Dice scores and sensitivity rates depending on the imaging modality and input dimension. Box edges mark the 25th and 75th percentiles, the central box line marks the median value.



Figure 4: Three exemplary patient cases, representing very high (upper row), average (middle row) and unsatisfactory (lower row) segmentation accuracies produced with 3D image input. Corresponding Dice scores are stated in the lower right corners. Green contours display the expertly annotated data as ground truth and blue contours represent our automatically produced segmentations. From left to right: (a) original  $T_1$ -weighted MRI sequence, (b) original  $T_2$ -weighted MRI sequence, (c) result with only  $T_1$ -weighted image data, (d) result with only  $T_2$ -weighted image data, (e) result with combined  $T_1$ - and  $T_2$ -weighted image data.

bone structures, which are additionally enhanced by the application of contrast agents. Multi-modal input, i.e. combining contrast-enhanced  $T_1$ -weighted images with  $T_2$ -weighted MR data showed only small

improvements regarding the mean accuracy and a somewhat reduced standard deviation (see Tab. 1). Applying solely  $T_2$ -weighted images yielded worse results, most likely due to the inferior image constrast between necrosis and surrounding tissues. Hence, they rather support and improve robustness in combination with the contrast-enhanced  $T_1$ -weighted input. With respect to the input dimension, it was found that on average 2D image input yielded higher scores than volumetric image input, which could be attributed to the higher number of trainable parameters (1,400,000 vs. 4,000,000) to be optimized and therefore, increased network complexity.

### 4 CONCLUSION

Automatic necrosis zone segmentation in followup MR scans after RF ablations of spinal metastases has the potential to quantify and objectify the treatment outcome validation. It provides important information regarding the improvement of ablation procedures and it may help understanding and predicting possible tumor reoccurrence. We proposed a CNNbased segmentation approach and examined the impact of various input modalities and dimensions on the segmentation accuracy. Our results were on par with those of Egger et al. (Egger et al., 2015), which were the only quantitative results available (77.2.0 % vs 77 %), altough the latter focused on necrotized liver lesions in CT imaging. Overall, our study indicates promising results and constitutes a valuale approach towards this ambitious and challenging issue.

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