Deep 3D Convolutional Encoder Networks with Shortcuts for Multiscale Feature Integration Applied to Multiple Sclerosis Lesion Segmentation

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Abstract—We propose a novel segmentation approach based on deep 3D convolutional encoder networks with shortcut connections and apply it to the segmentation of multiple sclerosis (MS) lesions in magnetic resonance images. Our model is a neural network that consists of two interconnected pathways, a convolutional pathway, which learns increasingly more abstract and higher-level image features, and a deconvolutional pathway, which predicts the final segmentation at the voxel level. The joint training of the feature extraction and prediction pathways allows for the automatic learning of features at different scales that are optimized for accuracy for any given combination of image types and segmentation task. In addition, shortcut connections between the two pathways allow high- and low-level features to be integrated, which enables the segmentation of lesions across a wide range of sizes. We have evaluated our method on two publicly available data sets (MICCAI 2008 and ISBI 2015 challenges) with the results showing that our method performs comparably to the top-ranked state-of-the-art methods, even when only relatively small data are available for training. In addition, we have evaluated our method on a large data set from an MS clinical trial, with a comparison of network architectures of different depths and with and without shortcut connections. The results show that increasing depth from three to seven layers improves performance, and adding shortcut connections further increases accuracy. For a direct comparison, we also ran the same data set through five freely available and widely used MS lesion segmentation methods, namely EMS, LST-LPA, LST-LGA, Lesion-TOADS, and SLS. The results show that our method consistently outperforms these other methods across a wide range of lesion sizes.

Index Terms—Segmentation, multiple sclerosis lesions, magnetic resonance imaging (MRI), deep learning, convolutional neural networks, machine learning

I. INTRODUCTION

MULTIPLE sclerosis (MS) is an inflammatory and demyelinating disease of the central nervous system with pathology that can be observed in vivo by magnetic resonance imaging (MRI). MS is characterized by the formation of lesions, primarily visible in the white matter on conventional MRI. Imaging biomarkers based on the delineation of lesions, such as lesion load and lesion count, have established their importance for assessing disease progression and treatment effect. However, lesions vary greatly in size, shape, intensity and location, which makes their automatic and accurate segmentation challenging.

Many automatic methods have been proposed for the segmentation of MS lesions over the last two decades [12], which can be classified into unsupervised and supervised methods. Unsupervised methods do not require a labeled data set for training. Instead, lesions are identified as an outlier of, e.g., a subject-specific generative model of tissue intensities [34], [41], [43], or a generative model of image patches representing a healthy population [44]. Alternatively, clustering methods have been used to segment healthy and lesion tissue, where lesions are modelled as a separate tissue class [35], [39]. In many methods, spatial priors of healthy tissues are used to reduce false positives. For example, in addition to modelling MS lesions as a separate intensity cluster, Lesion-TOADS [35] employs topological and statistical atlases to produce a topology-preserving segmentation of all brain tissues.

To account for local changes of the tissue intensity distributions, Tomas-Fernandez et al. [41] combined the subject-specific model of the global intensity distributions with a voxel-specific model calculated from a healthy population, where lesions are detected as outliers of the combined model. A major challenge of unsupervised methods is that outliers are often not specific to lesions and can also be caused by intensity inhomogeneities, partial volume, imaging artifacts, and small anatomical structures such as blood vessels, which leads to the generation of false positives. To overcome these limitations, Roura et al. [32] employed an additional set of rules to remove false positives, while Schmidt et al. [34] used a conservative threshold for the initial detection of lesions, which are later grown in a separate step to yield an accurate delineation.

Current supervised approaches typically start with a set of features, which can range from small and simple to large and highly variable, and are either predefined by the user [13], [14], [37] or gathered in a feature extraction step such as by deep learning [45]. Voxel-based segmentation algorithms [13], [45] feed the features and labels of each voxel into a
general classification algorithm, such as a random forest, to classify each voxel and to determine which set of features are the most important for segmentation in the particular domain. Voxel features and the labels of neighboring voxels can be incorporated into Markov random field-based (MRF-based) approaches to produce a spatially consistent segmentation. As a strategy to reduce false positives, Subbanna et al. combined a voxel-level MRF with a regional MRF, which integrates a large set of intensity and textural features extracted from the regions produced by the voxel-level MRF with the labels of neighboring nodes of the regional MRF. Library-based approaches leverage a library of pre-segmented images to carry out the segmentation. For example, Guizard et al. proposed a segmentation method based on an extension of the non-local means algorithms. The centers of patches at every voxel location are classified based on matched patches from a library containing pre-segmented images, where multiple matches are weighted using a similarity measure based on rotation-invariant features.

A recent breakthrough for the automatic segmentation using deep learning comes from the domain of cell membrane segmentation, in which Cireșan et al. proposed classifying the centers of image patches directly using a convolutional neural network (CNN) without a dedicated feature extraction step. Instead, features are learned indirectly within the lower layers of the neural network during training, while the higher layers can be regarded as performing the classification, which allows the learning of features that are specifically tuned to the segmentation task. However, the time required to train patch-based methods can make the approach infeasible when the size and number of patches are large.

Recently, different CNN architectures have been proposed that are able to feed through entire images, which removes the need to select representative patches, eliminates redundant calculations where patches overlap, and therefore these models scale up more efficiently with image resolution. Kang et al. introduced the fully convolutional neural network (fCNN) for the segmentation of crowds in surveillance videos. However, fCNNs produce segmentations of lower resolution than the input images due to the successive use of convolutional and pooling layers, both of which reduce the dimensionality. To predict segmentations of the same resolution as the input images, we recently proposed using a 3-layer convolutional encoder network (CEN) for MS lesion segmentation. The combination of convolutional and deconvolutional layers allows our network to produce segmentations that are of the same resolution as the input images.

Another limitation of the traditional CNN is the trade-off between localization accuracy, represented by lower-level features, and contextual information, provided by higher-level features. To overcome this limitation, Long et al. proposed fusing the segmentations produced by the lower layers of the network with the upsampled segmentations produced by higher layers. However, using only low-level features was not sufficient to produce a good segmentation at the lowest layers, which is why segmentation fusion was only performed for the three highest layers. Instead of combining the segmentations produced at different layers, Ronneberger et al. proposed combining the features of different layers to calculate the final segmentation directly at the lowest layer using an 11-layer u-shaped network architecture called u-net. Their network is composed of a traditional contracting path (first half of the u), but augmented with an expanding path (last half of the u), which replaces the pooling layers of the contracting path with upsampling operations. To leverage both high- and low-level features, shortcut connections are added between corresponding layers of the two paths. However, upsampling cannot fully compensate for the loss of resolution, and special handling of the border regions is still required.

We propose a new convolutional network architecture that combines the advantages of a CEN and a u-net. Our network is divided into two pathways, a traditional convolutional pathway, which consists of alternating convolutional and pooling layers, and a deconvolutional pathway, which consists of alternating deconvolutional and unpooling layers and predicts the final segmentation. Similar to the u-net, we introduce shortcut connections between layers of the two pathways. In contrast to the u-net, our network uses deconvolution instead of upsampling in the expanding pathway and predicts segmentations that have the same resolution as the input images and therefore does not require special handling of the border regions. We have evaluated our method on two widely used publicly available data sets for the evaluation of MS lesion segmentation methods and a large in-house data set from an MS clinical trial, with a comparison of network architectures of different depths and with and without shortcut connections. The results will be presented in detail in Section III.

II. Methods

In this paper, the task of segmenting MS lesions is defined as finding a function $s$ that maps multi-modal images $I$, e.g., $I = (I_{FLAIR}, I_T)$, to corresponding binary lesion masks $S$, where 1 denotes a lesion voxel and 0 denotes a non-lesion voxel. Given a set of training images $I_n$, $n \in \mathbb{N}$, and corresponding segmentations $S_n$, we model finding an appropriate function for segmenting MS lesions as an optimization problem of the following form

$$\hat{s} = \arg \min_{s \in S} \sum_n E(S_n, s(I_n)),$$

where $S$ is the set of possible segmentation functions, and $E$ is an error measure that calculates the dissimilarity between ground truth segmentations and predicted segmentations.

A. Model Architecture

The set of possible segmentation functions, $S$, is modeled by the convolutional encoder network with shortcut connections (CEN-s) illustrated in Fig. 1. A CEN-s is a type of convolutional neural network (CNN) that is divided into two interconnected pathways, the convolutional pathway and the deconvolutional pathway. The convolutional pathway...
consists of alternating convolutional and pooling layers. The input layer of the convolutional pathway is composed of the image voxels $x_i^{(0)}(\vec{p})$, $i \in [1, C]$, where $i$ indexes the modality or input channel, $C$ is the number of modalities or channels, and $\vec{p} \in \mathbb{N}^3$ are the coordinates of a particular voxel. The convolutional layers automatically learn a feature hierarchy from the input images. A convolutional layer is a deterministic function of the following form

$$x_j^{(l)} = \max \left( 0, \sum_{i=1}^C \tilde{w}_c^{(l)}_{ij} \ast x_i^{(l-1)} + b_j^{(l)} \right),$$

where $l$ is the index of a convolutional layer, $x_j^{(l)}$, $j \in [1, F]$, denotes the feature map corresponding to the trainable convolution filter $w_c^{(l)}_{ij}$, $F$ is the number of filters of the current layer, $b_j^{(l)}$ are trainable bias terms, $\ast$ denotes valid convolution, and $\tilde{w}$ denotes a flipped version of $w$, i.e., $\tilde{w}(\alpha) = w(-\alpha)$. To be consistent with the inference rules of convolutional restricted Boltzmann machines (convRBMs) [27], which are used for pre-training, convolutional layers convolve the input signal with flipped filter kernels, while deconvolutional layers calculate convolutions with non-flipped filter kernels. We use rectified linear units [30] in all layers except for the output layers, which have shown to improve the classification performance of CNNs [25]. A convolutional layer is followed by an average pooling layer [33] that halves the number of units in each dimension by calculating the average of each block of $2 \times 2 \times 2$ units per channel.

The deconvolutional pathway consists of alternating deconvolutional and unpooling layers with shortcut connections to the corresponding convolutional layers. The first deconvolutional layer uses the extracted features of the convolutional pathway to calculate abstract segmentation features

$$y_i^{(L-1)} = \max \left( 0, \sum_{j=1}^F w_{d,ij}^{(L)} \ast y_j^{(L)} + c_i^{(L-1)} \right),$$

where $y_i^{(L)} = x_i^{(L)}$, $L$ denotes the number of layers of the convolutional pathway, $w_{d,ij}^{(L)}$ and $c_i^{(L-1)}$ are trainable parameters of the deconvolutional layer, and $\ast$ denotes full convolution. To be consistent with the general notation of deconvolutions [47], the non-flipped version of $w$ is convolved with the input signal.

Subsequent deconvolutional layers use the activations of the previous layer and corresponding convolutional layer to calculate more localized segmentation features

$$y_i^{(l)} = \max \left( 0, \sum_{j=1}^F w_{d,ij}^{(l+1)} \ast y_j^{(l+1)} + \sum_{j=1}^F w_{s,ij}^{(l+1)} \ast x_j^{(l+1)} + c_i^{(l)} \right),$$

where $l$ is the index of a deconvolutional layer with shortcut, and $w_{s,ij}^{(l+1)}$ are the shortcut filter kernels connecting the activations of the convolutional pathway with the activations of the deconvolutional pathway. The last deconvolutional layer integrates the low-level features extracted by the first convolutional layer with the high-level features from the previous layer to calculate a probabilistic lesion mask

$$y_i^{(0)} = \text{sigmoid} \left( \sum_{j=1}^F \left( w_{d,1,ij}^{(1)} \ast y_j^{(1)} + w_{s,1,ij}^{(1)} \ast x_j^{(1)} \right) + c_i^{(0)} \right),$$

where we use the sigmoid function defined as $\text{sigmoid}(z) = \frac{1 + \exp(z)}{1 - \exp(-z)}$, $z \in \mathbb{R}$ instead of the rectified linear function in order to obtain a probabilistic segmentation with values in the range between 0 and 1. To produce a binary lesion mask from the probabilistic output of our model, we chose a fixed threshold such that the mean Dice similarity coefficient [10] is maximized on the training set and used the same threshold for the evaluation on the test set.
B. Gradient Calculation

The parameters of the model can be efficiently learned by minimizing the error $E$ for each sample of the training set, which requires the calculation of the gradient of $E$ with respect to the model parameters \[26\]. Typically, neural networks are trained by minimizing the sum of squared differences (SSD), which can be calculated for a single image as follows

$$E = \frac{1}{2} \sum_{\tilde{p}} \left( S(\tilde{p}) - y^{(0)}(\tilde{p}) \right)^2,$$

where $\tilde{p} \in \mathbb{N}^3$ are the coordinates of a particular voxel. The partial derivatives of the error with respect to the model parameters can be calculated using the delta rule and are given by

$$\frac{\partial E}{\partial u_{d,i,j}} = \delta_{d,i}^{(l-1)} \ast y_j^{(l)}$$

$$\frac{\partial E}{\partial u_{c,i,j}} = \delta_{c,i}^{(l)} \ast x_i^{(l-1)}$$

$$\frac{\partial E}{\partial \theta_{c,i}} = x_i^{(l-1)} \ast c_{c,j}^{(l)}$$

For the first layer, $\delta_{d,1}^{(0)}$ can be calculated by

$$\delta_{d,1}^{(0)} = (y_1^{(0)} - S)y_1^{(0)}(1 - y_1^{(0)}).$$

The derivatives of the error with respect to the parameters of the other layers can be calculated by applying the chain rule of partial derivatives, which yields to

$$\delta_{d,1}^{(l)} = (\tilde{w}_{d,i,j}^{(l-1)} + \delta_{d,i}^{(l-1)})I(y_j^{(l)} > 0),$$

$$\delta_{c,i}^{(l)} = (\tilde{w}_{c,i,j}^{(l)} + \delta_{c,j}^{(l-1)})I(x_i^{(l)} > 0),$$

where $l$ is the index of a deconvolutional or convolutional layer, $\delta_{d,1}^{(l)} = \delta_{d,1}^{(l)}$, and $I(z)$ denotes the indicator function defined as 1 if the predicate $z$ is true and 0 otherwise. If a layer is connected through a shortcut, $\delta_{c,j}^{(l)}$ needs to be adjusted by propagating the error back through the shortcut connection. In this case, $\delta_{c,j}^{(l)}$ is calculated by

$$\delta_{c,j}^{(l)} = (\tilde{w}_{c,j}^{(l)} + \delta_{c,j}^{(l-1)})I(x_i^{(l)} > 0),$$

where $\tilde{w}_{c,j}^{(l)}$ denotes the activation of unit $\delta_{c,j}^{(l)}$ before taking the shortcut connection into account.

The sum of squared differences is a good measure of classification accuracy, if the two classes are fairly balanced. However, if one class contains vastly more samples, as is the case for lesion segmentation, the error measure is dominated by the majority class and consequently, the neural network would learn to ignore the minority class. To overcome this problem, we use a combination of sensitivity and specificity, which can be used together to measure classification performance even for vastly unbalanced problems. More precisely, the final error measure is a weighted sum of the mean squared difference of the lesion voxels (sensitivity) and non-lesion voxels (specificity), reformulated to be error terms:

$$E = \frac{r \sum_{\tilde{p}} (S(\tilde{p}) - y^{(0)}(\tilde{p}))^2}{\sum_{\tilde{p}} S(\tilde{p})} + \frac{(1 - r) \sum_{\tilde{p}} (S(\tilde{p}) - y^{(0)}(\tilde{p}))^2 (1 - S(\tilde{p}))}{\sum_{\tilde{p}} (1 - S(\tilde{p}))}.$$

We formulate the sensitivity and specificity errors as squared errors in order to yield smooth gradients, which makes the optimization more robust. The sensitivity ratio $r$ can be used to assign different weights to the two terms. Due to the large number of non-lesion voxels, weighting the specificity error higher is important, but based on preliminary experimental results \[4\], the algorithm is stable with respect to changes in $r$, which largely affects the threshold used to binarize the probabilistic output. A detailed evaluation of the impact of the sensitivity ratio on the learned model is presented in Section III-D.

To train our model, we must compute the derivatives of the modified objective function with respect to the model parameters. Equations \[7\] and \[14\] are a consequence of the chain rule and independent of the chosen similarity measure. Hence, we only need to derive the update rule for $\delta_{d,1}^{(0)}$. With $\alpha = 2r(\sum_{\tilde{p}} S(\tilde{p}))^{-1}$ and $\beta = 2(1 - r)(\sum_{\tilde{p}} (1 - S(\tilde{p})))^{-1}$, we can rewrite $E$ as

$$E = \frac{1}{2} \sum_{\tilde{p}} (\alpha S(\tilde{p}) + \beta (1 - S(\tilde{p}))) (S(\tilde{p}) - y^{(0)}(\tilde{p}))^2.$$

Our objective function is similar to the SSD, with an additional multiplicative term applied to the squared differences. The additional factor is constant with respect to the model parameters. Consequently, $\delta_{d,1}^{(0)}$ can be derived analogously to the SSD case, and the new factor is simply carried over:

$$\delta_{d,1}^{(0)} = (\alpha S + \beta (1 - S))(y_1^{(0)} - S)y_1^{(0)}(1 - y_1^{(0)}).$$

C. Training

At the beginning of the training procedure, the model parameters need to be initialized and the choice of the initial parameters can have a big impact on the learned model \[40\]. In our experiments, we found that initializing the model using pre-training \[16\] on the input images was required in order to be able to fine-tune the model using the ground truth segmentations without getting stuck early in a local minimum. Pre-training can be performed layer by layer \[15\] using a stack of convRBMs (see Fig. \[4\]), thereby avoiding the potential problem of vanishing or exploding gradients \[17\]. The first convRBM is trained on the input images, while subsequent convRBMs are trained on the hidden activations of the previous convRBM. After all convRBMs have been trained, the model parameters of the CEN-s can be initialized as follows (showing the first convolutional and the last deconvolutional layers only, see Fig. \[10\])

$$w^{(1)}_c = \tilde{w}^{(1)}, \quad w^{(1)}_d = 0.5 \tilde{w}^{(1)}$$

$$b^{(1)} = \tilde{b}^{(1)}, \quad c^{(0)} = \tilde{c}^{(1)},$$

\[17\]

\[18\]
where \( \hat{w}^{(1)} \) are the filter weights, \( \hat{b}^{(1)} \) are the hidden bias terms, and \( \hat{c}^{(1)} \) are the visible bias terms of the first convRBM.

A major challenge for gradient-based optimization methods is the choice of an appropriate learning rate. Classic stochastic gradient descent \( \text{SGD} \) uses a fixed or decaying learning rate, which is the same for all parameters of the model. However, the partial derivatives of parameters of different layers can vary substantially in magnitude, which can require different learning rates. In recent years, there has been an increasing interest in developing methods for automatically choosing independent learning rates. Most methods (e.g., AdaGrad \[11\], AdaDelta \[16\], RMSprop \[9\], and Adam \[24\]) collect different statistics of the partial derivatives over multiple iterations and use this information to set an adaptive learning rate for each parameter. This is especially important for the training of deep networks, where the optimal learning rates often differ greatly for each layer. In our initial experiments, networks obtained by training with AdaDelta, RMSprop, and Adam performed comparably well, but AdaDelta was the most robust to the choice of hyperparameters, so we used AdaDelta for all results reported.

D. Implementation

Pre-training and fine-tuning were performed using a highly optimized GPU-accelerated implementation of 3D convRBM and CENs that performs training in the frequency domain \[3\]. Our frequency domain implementation significantly speeds up the training by mapping the calculation of convolutions to simple element-wise multiplications, while adding only a small number of Fourier transforms. This is especially beneficial for the training on 3D volumes, due to the increased number of weights of 3D kernels compared to 2D. Although GPU-accelerated deep learning libraries based on cuDNN \[5\] are publicly available (e.g., \[1\], \[7\], \[21\]), we trained our models using our own implementation because we have found that it performs the most computationally intensive training operations 6 times faster than cuDNN in a direct comparison.

III. EXPERIMENTS AND RESULTS

We evaluated our method on two publicly available data sets (from the MICCAI 2008 and ISBI 2015 lesion segmentation challenges), which allows for a direct comparison with many state-of-the-art methods. In addition, we have used a much larger data set containing four different MRI sequences from a multi-center clinical trial in relapsing remitting MS, which tends to have the most heterogeneity among the MS subtypes. This data test is challenging due to the large variability in lesion size, shape, location, and intensity as well as varying contrasts produced by different scanners. The clinical trial data set was used to carry out a detailed analysis of different CEN architectures using different combinations of modalities, with a comparison to five publicly available state-of-the-art methods.

A. Data Sets and Pre-processing

1) Public data sets: The data set of the MICCAI 2008 MS lesion segmentation challenge \[36\] consists of 43 T1-weighted (T1w), T2-weighted (T2w), and FLAIR MRIs, divided into 20 training cases for which ground truth segmentations are made publicly available, and 23 test cases. After training the model on the 20 training cases, we used the trained model to segment the 23 test cases, which were sent to the challenge organizers for independent evaluation.

The data set of the ISBI 2015 longitudinal MS lesion segmentation challenge consists of 21 visit sets, each with T1w, T2w, proton density-weighted (PDw), and FLAIR MRIs. The challenge was not open for new submissions at the time of writing this article. Therefore, we evaluated our method on the training set using leave-one-subject-out cross-validation, following the evaluation protocol of the second \[20\] and third place \[29\] methods from the challenge proceedings. The paper of the first place method does not have sufficient details to replicate their evaluation.

2) Clinical trial data set: The data set was collected from 67 different scanning sites using different 1.5T and 3T scanners, and consists of T1w, T2w, PDw, and FLAIR MRIs from 195 subjects, most with two time points (377 visit sets in total). The image dimensions and voxel sizes vary by site, but most of the T1w images have close to 1 mm isotropic voxels, while the other images have voxel sizes close 1 mm x 1 mm x 3 mm. All images were skull-stripped using the brain extraction tool (BET) \[18\], followed by an intensity normalization to the interval \([0, 1]\), and a 6 degree-of-freedom intra-subject registration using one of the 3 mm scans as the target image to align the different modalities. To speed-up the training, all images were cropped to a 164 x 206 x 52 voxel subvolume with the brain roughly centered. The ground truth segmentations were produced using an existing semi-automatic 2D region-growing technique, which has been used successfully in a number of large MS clinical trials (e.g., \[23\], \[42\]). Each lesion was manually identified by an experienced radiologist and then interactively grown from the seed point by a trained technician.

We divided the data set into a training \((n = 250)\), validation \((n = 50)\), and test set \((n = 77)\) such that images of each set were acquired from different scanning sites. The training, validation, and test sets were used for training our models, for monitoring the training progress, and to evaluate performance, respectively. The training set was also used to perform parameter tuning of the other methods used for comparison. Pre-training and fine-tuning of our 7-layer CENs took approximately 27 hours and 37 hours, respectively, on a single GeForce GTX 780 graphics card. Once the network is trained, new multi-contrast images can be segmented in less than one second.

B. Comparison to Other Methods

We compared our method with the five publicly available methods, some of which are widely used for clinical research and are established in the literature (e.g., \[14\], \[37\], \[39\]) as reference points for comparison. These five methods include:

1) Expectation maximization segmentation (EMS) method \[43\]:
2) Lesion growth algorithm (LST-LGA) [34], as implemented in the Lesion Segmentation Toolbox (LST) version 2.0.11;
3) Lesion prediction algorithm (LST-LPA) also implemented in the same LST toolbox;
4) Lesion-TOADS version 1.9 R [35]; and
5) Salem Lesion Segmentation (SLS) toolbox [32].

The Lesion-TOADS software only takes T1w and FLAIR MRIs and has no tunable parameters, so we used the default parameters to carry out the segmentations. The performance of EMS depends on the choice of the Mahalanobis distance $\kappa$, the threshold $t$ used to binarize the probabilistic segmentation, and the modalities used. We applied EMS to segment lesions using two combinations of modalities: a) T1w, T2w, and PDw, as used in the original paper [43], and b) all four available modalities (T1w, T2w, PD2, FLAIR). We compared the segmentations produced for all combinations of $\kappa = 2.0, 2.2, \ldots, 4.6$ and $t = 0.05, 0.10, \ldots, 1.00$ with the ground truth segmentations on the training set and chose the values that maximized the average DSC ($\kappa = 2.6$, $t = 0.75$ for three modalities; $\kappa = 2.8$, $t = 0.9$ for four modalities).

The LST-LGA and LST-LPA of the LST toolbox only take T1w and FLAIR MRIs as input, and we used those modalities to tune the initial threshold $\kappa$ of LST-LGA for $\kappa = 0.05, 0.10, \ldots, 1.00$ and the threshold $t$ used by LST-LPA to binarize the probabilistic segmentations for $t = 0.05, 0.10, \ldots, 1.00$. The optimal parameters were $\kappa = 0.10$ and $t = 0.45$, respectively.

Similarly, the SLS toolbox, which is the most recently published work [32] also only takes T1w and FLAIR MRIs as input. This method uses an initial brain tissue segmentation obtained from the T1w images and segments lesions by treating lesional pixels as outliers to the normal appearing GM tissue on the FLAIR images. This method has three key parameters: $\omega_{\text{GM}}$, $\omega_{\text{CSF}}$, and $\omega_{\text{LS}}$. We tuned these parameters on the training set via a grid-search over a range of values as suggested in [12].

C. Measures of Segmentation Accuracy

We used the following four measures to produce a comprehensive evaluation of segmentation accuracy as there is generally no single measure that is sufficient to capture all information relevant to the quality of a produced segmentation [12].

The first measure is the Dice similarity coefficient (DSC) [10] that computes a normalized overlap value between the produced and ground truth segmentations, and is defined as

$$\text{DSC} = \frac{2 \times TP}{2 \times TP + FP + FN},$$

where $TP$, $FP$, and $FN$ denote the number of true positive, false positive, and false negative voxels, respectively. A value of 100% indicates a perfect overlap of the produced segmentation and the ground truth. The DSC incorporates measures of overlap and underestimation into a single metric, which makes it a suitable measure to compare overall segmentation accuracy.

We also measured the relative absolute volume difference between the ground truth and the produced segmentation by computing their volumes ($\text{Vol}$), i.e.,

$$\text{VD} = \frac{\text{Vol}(\text{Seg}) - \text{Vol}(\text{GT})}{\text{Vol}(\text{GT})},$$

where $\text{Seg}$ and $\text{GT}$ denote the obtained segmentation and ground truth, respectively. However, it has been noted [12] that wide variability exists even between the lesion segmentations of trained experts, and thus, the achieved volume differences reported in the literature have ranged from 10% to 68%.

For more precise evaluation, we have also included the lesion-wise true positive rate (LTPR) and the lesion-wise false positive rate (LFPR) that are much more sensitive in measuring the segmentation accuracy of smaller lesions, which are important to detect when performing early disease diagnosis [12]. More specifically, the LTPR measures the true positive rate (TPR) on a per lesion-basis and is defined as

$$\text{LTPR} = \frac{\text{LTP}}{\#\text{RL}},$$

where LTP denotes the number of lesion true positive, i.e., the number lesions in the reference segmentation that overlap with a lesion in the produced segmentation, and $\#\text{RL}$ denotes the total number of lesions in the reference segmentation. An LTPR with a value of 100% indicates that all lesions are correctly identified. Similarly, the lesion-wise false positive rate (FPR) measures the fraction of the segmented lesions that are not in the ground truth and is defined as

$$\text{LFPR} = \frac{\text{LFP}}{\#\text{PL}},$$

where LFP denotes the number of lesion false positives, i.e., the number of lesions in the produced segmentation that do not overlap with a lesion in the reference segmentation, and $\#\text{PL}$ denotes the total number of lesions in the produced segmentation. An LFPR with a value of 0% indicates that no lesions were incorrectly identified.

D. Training Parameters

The most influential parameters of the training method are the number of epochs and the sensitivity ratio. Fig. 2 shows the mean DSC evaluated on the training and validation sets of the clinical data set as computed during training of a 7-layer CEN-s up to 500 epochs. The mean DSC scores increase monotonically, but the improvements are minor after 400 epochs. The optimal number of epochs is a trade-off between accuracy and time required for training. Due to the relatively small improvements after 400 epochs, we decided to stop the training procedure at 500 epochs. For the challenge data sets, due to their small sizes, we did not employ a subset of the data for a dedicated validation set to choose the number of epochs. Instead, we set the number of epochs to 2500, which corresponds to roughly the same number of gradient updates compared to the clinical trial data set.

To determine an effective sensitivity ratio, we measured the performance on the validation set over a range of values. For each choice of ratio, we binarized the segmentations using a
threshold that maximized the DSC on the training set. Fig. 3 shows a set of ROC curves for different choices of the sensitivity ratio ranging from 0.01 to 0.10 and the corresponding optimal thresholds. The plots illustrate our findings that our method is not sensitive to the choice of the sensitivity ratio, which mostly affects the optimal threshold. We chose a fixed sensitivity ratio of 0.02 for all our experiments.

### E. Comparison on Public Data Sets

To allow for a direct comparison with a large number of state-of-the-art methods, we evaluated our method on the MICCAI 2008 MS lesion segmentation challenge [36] and the ISBI 2015 longitudinal MS lesion segmentation challenge. We have previously shown that approximately 100 images are required to train the 3-layer CEN without overfitting [4] and we expect the required number of images to be even higher when adding more layers. Due to the relatively small size of the training data sets provided by the two challenges, we used a CEN with only three layers on these data sets to reduce the risk of overfitting. The parameters of the models are summarized in Table I.

A comparison of our method with other state-of-the-art methods evaluated on the MICCAI challenge test data set is summarized in Table II. Our method ranked 6th (2nd if only considering methods with only one submission, i.e., without subsequent parameter-tuning and adjustments) out of 52 entries submitted to the challenge, outperforming the recent SLS by Roura et al. [32], and popular methods such as the random forest approach by Geremia et al. [13], and Lesion-TOADS by Shiee et al. [35], but not as well as the patch-based segmentation approach by Guizard et al. [14], or the MOPS approach by Tomas-Fernandez et al. [41], which used additional images to build the intensity model of a healthy population. This is a very promising result for the first submission of our method given the simplicity of the model and the small training set size.

In addition, we evaluated our method on the 21 publicly available labeled cases from the ISBI 2015 longitudinal MS lesion segmentation challenge. The challenge organizers have only released the names of the top three teams, only two of which have published a summary of their mean DSC, LTPR, and LFPR scores for both raters to allow for a direct comparison. Following the evaluation protocol of the second [20] and third [29] place methods, we trained our model using leave-one-subject-out cross-validation on the training images and compared our results to the segmentations provided by both raters. Table III summarizes the performance of our method, the two other methods for comparison, and the performance of the two raters when compared against each other. Compared to the second and third place methods, our method was more sensitive and produced significantly higher LTPR scores, but also had more false positives, which resulted in slightly lower but still comparable DSC scores. This is again a promising result on a public data set.
TABLE III
COMPARISON OF OUR METHOD WITH THE SECOND AND THIRD RANKED METHODS FROM THE ISBI MS LESION SEGMENTATION CHALLENGE.

<table>
<thead>
<tr>
<th>Method</th>
<th>Rater 1 DSC</th>
<th>Rater 1 LTPR</th>
<th>Rater 1 LFPR</th>
<th>Rater 2 DSC</th>
<th>Rater 2 LTPR</th>
<th>Rater 2 LFPR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maier et al. [29]</td>
<td>70.4</td>
<td>61.1</td>
<td>13.5</td>
<td>68.1</td>
<td>50.1</td>
<td>12.7</td>
</tr>
<tr>
<td>Maier et al. (GT2)</td>
<td>70</td>
<td>53</td>
<td>48</td>
<td>65</td>
<td>37</td>
<td>44</td>
</tr>
<tr>
<td>Our method (GT1)</td>
<td>68.4</td>
<td>74.5</td>
<td>54.5</td>
<td>64.4</td>
<td>63.0</td>
<td>52.8</td>
</tr>
<tr>
<td>Our method (GT2)</td>
<td>68.3</td>
<td>78.3</td>
<td>64.5</td>
<td>65.8</td>
<td>69.3</td>
<td>61.9</td>
</tr>
</tbody>
</table>

Note: The evaluation was performed on the training set using leave-one-subject-out cross-validation. GT1 and GT2 denote that the model was trained with the segmentations provided by the first and second rater as the ground truth, respectively.

F. Comparison of Network Architectures, Input Modalities, and Publicly Available Methods on Clinical Trial Data

To determine the effect of network architecture, we compared the segmentation performance of three different networks using T1w and FLAIR MRIs. Speciﬁcally, we trained a 3-layer CEN and two 7-layer CENs, one with shortcut connections and one without. To investigate the effect of different input image types, we additionally trained two 7-layer CEN-s on the modalities used by EMS (T1w, T2w, PDw) and all four modalities (T1w, T2w, PDw, FLAIR). The parameters of the networks are given in Table IV and Table V. To roughly compensate for the anisotropic voxel size of the input images, we chose an anisotropic filter size of $9 \times 9 \times 5$.

In addition, we ran the five competing methods discussed in Section III-B with Lesion-TOADS, SLS, and the two LST methods using the T1w and FLAIR images, and EMS using three (T1w, T2w, PDw) and all four modalities in separate tests. A comparison of the segmentation accuracy of the trained networks and competing methods is summarized in Table VI.

All CEN architectures performed signiﬁcantly better than all other methods regardless of the input modalities, with LST-LGA being the closest in overall segmentation accuracy. Comparing CEN to LST-LGA, the improvements in the mean DSC scores ranged from 3 percentage points (pp) for the 3-layer CEN to 17 pp for the 7-layer CEN with shortcut trained on all four modalities. The improved segmentation performance was mostly due to an increase in lesion sensitivity. LST-LGA achieved a mean lesion TPR of 37.50%, compared to 54.55% produced by the CEN with shortcut when trained on the same modalities, and 62.49% when trained on all four modalities, while achieving a comparable number of lesion false positives. The mean lesion FPRs and mean volume differences of LST-LGA and the 7-layer CEN-s were very close, when trained on the same modalities, and the CEN-s further reduced its FPR when trained on more modalities.

This experiment also showed that increasing the depth of the CEN and adding the shortcut connections both improve the segmentation accuracy. Increasing the depth of the CEN from three layers to seven layers improved the mean DSC by 3 pp. The improvement was conﬁrmed to be statistically signiﬁcant using a one-sided paired t-test ($p$-value of $1.3 \times 10^{-5}$). Adding a shortcut to the network further improved the segmentation accuracy as measured by the DSC by 3 pp. A second one-sided paired t-test was performed to confirm the statistical signiﬁcance of the improvement with a $p$-value of less than $1 \times 10^{-10}$.

The impact of increasing the depth of the network on the segmentation performance of very large lesions is illustrated in Fig. 4, where the true positive, false negative, and false positive voxels are highlighted in green, yellow, and red, respectively. The receptive ﬁeld of the 3-layer CEN has a size of only $17 \times 17 \times 9$ voxels, which reduces its ability to identify very large lesions marked by two white circles.

TABLE IV
PARAMETERS OF THE 3-LAYER CEN USED ON THE CLINICAL TRIAL DATA SET.

<table>
<thead>
<tr>
<th>Layer type</th>
<th>Kernel Size</th>
<th>#Filters</th>
<th>Image Size</th>
</tr>
</thead>
<tbody>
<tr>
<td>Convolutional</td>
<td>$9 \times 9 \times 5 \times 2$</td>
<td>32</td>
<td>$164 \times 206 \times 52 \times 2$</td>
</tr>
<tr>
<td>Deconvolutional</td>
<td>$9 \times 9 \times 5 \times 32$</td>
<td>1</td>
<td>$164 \times 206 \times 52 \times 1$</td>
</tr>
</tbody>
</table>

TABLE V
PARAMETERS OF THE 7-LAYER CEN-S USED ON THE CLINICAL TRIAL DATA SET.

<table>
<thead>
<tr>
<th>Layer type</th>
<th>Kernel Size</th>
<th>#Filters</th>
<th>Image Size</th>
</tr>
</thead>
<tbody>
<tr>
<td>Convolutional</td>
<td>$9 \times 9 \times 5 \times 2$</td>
<td>32</td>
<td>$156 \times 198 \times 48 \times 32$</td>
</tr>
<tr>
<td>Average Pooling</td>
<td>$2 \times 2 \times 2$</td>
<td>—</td>
<td>$78 \times 99 \times 24 \times 32$</td>
</tr>
<tr>
<td>Convolutional</td>
<td>$9 \times 10 \times 5 \times 32$</td>
<td>32</td>
<td>$70 \times 90 \times 20 \times 32$</td>
</tr>
<tr>
<td>Deconvolutional</td>
<td>$9 \times 10 \times 5 \times 32$</td>
<td>32</td>
<td>$78 \times 99 \times 24 \times 32$</td>
</tr>
<tr>
<td>Unpooling</td>
<td>$2 \times 2 \times 2$</td>
<td>—</td>
<td>$156 \times 198 \times 48 \times 32$</td>
</tr>
</tbody>
</table>

TABLE VI
COMPARISON OF THE SEGMENTATION ACCURACY OF DIFFERENT CEN MODELS, OTHER METHODS, AND INPUT MODALITIES.

<table>
<thead>
<tr>
<th>Method</th>
<th>DSC [%]</th>
<th>LTPR [%]</th>
<th>LFPR [%]</th>
<th>VD [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Input modalities: T1w and FLAIR</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3-layer CEN [4]</td>
<td>49.24</td>
<td>57.33</td>
<td>61.39</td>
<td>43.45</td>
</tr>
<tr>
<td>7-layer CEN</td>
<td>52.07</td>
<td>43.88</td>
<td>29.06</td>
<td>37.01</td>
</tr>
<tr>
<td>7-layer CEN-s</td>
<td>55.76</td>
<td>54.55</td>
<td>38.64</td>
<td>36.30</td>
</tr>
<tr>
<td>Lesion-TOADS [35]</td>
<td>40.04</td>
<td>56.56</td>
<td>82.90</td>
<td>49.36</td>
</tr>
<tr>
<td>SLS [52]</td>
<td>43.20</td>
<td>56.80</td>
<td>50.80</td>
<td>12.30</td>
</tr>
<tr>
<td>LST-LGA [34]</td>
<td>46.64</td>
<td>37.50</td>
<td>38.06</td>
<td>36.77</td>
</tr>
<tr>
<td>LST-LPA [39]</td>
<td>46.07</td>
<td>48.02</td>
<td>52.94</td>
<td>41.62</td>
</tr>
</tbody>
</table>

| Input modalities: T1w, T2w, and PDw |
| 7-layer CEN-s   | 61.18   | 52.00    | 36.68    | 29.38 |
| EMS [43]        | 42.94   | 44.80    | 76.58    | 49.29 |

Note: The table shows the mean of the Dice similarity coefﬁcient (DSC), lesion true positive rate (LTPR), and lesion false positive rate (LFPR). Because the volume difference (VD) is not limited to the interval $[0, 100]$, a single outlier can heavily affect the calculation of the mean. We therefore excluded outliers before calculating the mean of the VD for all methods using the box plot criterion.
In contrast, the 7-layer CEN has a receptive field size of 49 × 53 × 26 voxels, which allows it to learn features that can capture much larger lesions. Consequently, the 7-layer CEN, with and without shortcut, is able to learn a feature set that captures large lesions much better than the 3-layer CEN, which results in an improved segmentation. However, increasing the depth of the network without adding shortcut connections reduces the network’s sensitivity to very small lesions as illustrated in Fig. 5. In this example, the 3-layer CEN was able to detect three small lesions, indicated by the white circles, which were missed by the 7-layer CEN. Adding shortcut connections enables our model to learn a feature set that spans a wider range of lesion sizes, which increases the sensitivity to small lesions and, hence, allows the 7-layer CEN-s to detect all three small lesions (highlighted by the white circles), while still being able to segment large lesions.

G. Comparison for Different Lesion Sizes

To examine the effect of lesion size on segmentation performance, we stratified the test set into five groups based on their mean reference lesion size as summarized in Table VII. A comparison of segmentation accuracy and lesion detection measures of a 7-layer CEN-s trained on different input modalities and the best performing competing method LST-LGA for different lesion sizes is illustrated in Fig. 6. The 7-layer CEN-s outperformed LST-LGA for all lesion sizes except for very large lesions when trained on T1w and FLAIR MRIs. The advantage extended to all lesion sizes when the CEN-s was trained on all four modalities, which could not be done for LST-LGA. The differences were larger for smaller lesions, which are generally more challenging to segment for all methods. The differences between the two approaches were due to a higher sensitivity to lesions as measured by the LTPR, especially for smaller lesions, while the number of false positives was approximately the same for all lesion sizes.

IV. Discussion

The automatic segmentation of MS lesions is a very challenging task due to the large variability in lesion size, shape, intensity, and location, as well as the large variability of imaging contrasts produced by different scanners used in multicenter studies. Most unsupervised methods model lesions as an outlier class or a separate cluster in a subject-specific model, which makes them inherently robust to inter-subject and inter-scanner variability. However, outliers are often not specific to lesions and can also be caused by intensity inhomogeneities, partial volume, imaging artifacts, and small anatomical structures such as blood vessels, which leads to the generation of false positives. On the other hand, supervised methods can learn to discriminate between lesion and non-lesion tissue, but are more sensitive to the variability in lesion appearance and different contrasts produced by different scanners. To overcome these challenges, supervised methods require large data sets that span the variability in lesion appearance and careful pre-processing to match the imaging contrast of new images with those of the training set. Library-based approaches have shown great promise for the segmentation of MS lesions, but do not scale well to very large data sets due to the large amount of memory required to store comprehensive sample libraries and the time required to scan such libraries for matching patches. On the other hand, parametric deep learning models such as convolutional neural networks scale much better to large training sets, because the size required to store the model is independent of the training set size, and the operations required for training and inference are inherently parallelizable, which allows them to take advantage of very fast GPU-accelerated computing hardware. Furthermore, the combination of many nonlinear processing units allows them to learn features that are robust under large variability, which is crucial for the segmentation of MS lesions.
Convolutional neural networks were originally designed to classify entire images and designing networks that can segment images remains an important research topic. Early approaches have formulated the segmentation problem as a patch-wise classification problem, which allows them to directly use established classification network architectures for image segmentation. However, a major limitation of patch-based deep learning approaches is the time required for training and inference. Fully convolutional networks can perform the segmentation much more efficiently, but generally lack the precision to perform voxel-accurate segmentation and cannot handle unbalanced classes.

To overcome these challenges, we have presented a new method for the automatic segmentation of MS lesions based on deep convolutional encoder networks with shortcut connections. The joint training of the feature extraction and prediction pathways allows for the automatic learning of features at different scales that are tuned for a given combination of image types and segmentation task. Shortcuts between the two pathways allow high- and low-level features to be leveraged at the same time for more consistent performance across scales.

In addition, we have proposed a new objective function based on the combination of sensitivity and specificity, which makes the objective function inherently robust to unbalanced classes such as MS lesions, which typically comprise less than 1% of all image voxels. We have evaluated our method on two publicly available data sets and a large data set from an MS clinical trial, with the results showing that our method performs comparably to the best state-of-the-art methods, even for relatively small training set sizes. We have also shown that when a suitably large training set is available, our method is able to segment MS more accurately than widely-used competing methods such as EMS, LST-LGA, SLS, and Lesion-TOADS. The substantial gains in accuracy were mostly due to an increase in lesion sensitivity, especially for small lesions. Overall, our proposed CEN with shortcut connections performed consistently well over a wide range of lesion sizes.

Our segmentation framework is very flexible and can be easily extended. One such extension could be to incorporate prior knowledge about the tissue type of each non-lesion voxel into the segmentation procedure. The probabilities of each tissue class could be precomputed by a standard segmentation method, after which they can be added as an additional channel to the input units of the CEN, which would allow the CEN to take advantage of intensity information from different modalities and prior knowledge about each tissue class to carry out the segmentation. In addition, our method can be applied to other segmentation tasks. Although we have only focused on the segmentation of MS lesions in this paper, our method does not make any assumptions specific to MS lesion segmentation. The features required to carry out the segmentation are solely learned from training data, which allows our method to be used to segment different types of pathology or anatomy when a suitable training set is available.

Acknowledgements

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References


