



# BRAVE-NET: Fully Automated Arterial Brain Vessel Segmentation in Patients With Cerebrovascular Disease

Adam Hilbert<sup>1\*</sup>, Vince I. Madai<sup>1,2</sup>, Ela M. Akay<sup>1</sup>, Orhun U. Aydin<sup>1</sup>, Jonas Behland<sup>1</sup>, Jan Sobesky<sup>3,4</sup>, Ivana Galinovic<sup>3</sup>, Ahmed A. Khalil<sup>3,5,6,7</sup>, Abdel A. Taha<sup>8</sup>, Jens Wuerfel<sup>9</sup>, Petr Dusek<sup>10</sup>, Thoralf Niendorf<sup>11</sup>, Jochen B. Fiebach<sup>3</sup>, Dietmar Frey<sup>1</sup> and Michelle Livne<sup>1</sup>

<sup>1</sup> CLAIM - Charité Lab for Artificial Intelligence in Medicine, Charité Universitätsmedizin Berlin, Berlin, Germany, <sup>2</sup> Faculty of Computing, Engineering and the Built Environment, School of Computing and Digital Technology, Birmingham City University, Birmingham, United Kingdom, <sup>3</sup> Centre for Stroke Research Berlin, Charité Universitätsmedizin Berlin, Berlin, Germany, <sup>4</sup> Johanna-Etienne-Hospital, Neuss, Germany, <sup>5</sup> Department of Neurology, Max Planck Institute for Human Cognitive and Brain Sciences, Leipzig, Germany, <sup>6</sup> Berlin School of Mind and Brain, Mind, Brain, Body Institute, Humboldt-Universität Berlin, Berlin, Germany, <sup>7</sup> Biomedical Innovation Academy, Berlin Institute of Health, Berlin, Germany, <sup>8</sup> Research Studio Data Science, Research Studios Austria, Salzburg, Austria, <sup>9</sup> Department Biomedical Engineering, Medical Image Analysis Center AG, University of Basel, Basel, Switzerland, <sup>10</sup> Department of Neurology, 1st Faculty of Medicine, Centre of Clinical Neuroscience, General University Hospital in Prague, Charles University, Prague, Czechia, <sup>11</sup> Berlin Ultrahigh Field Facility, Max Delbrück Center for Molecular Medicine in the Helmholtz Association, Berlin, Germany

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### \*Correspondence:

Adam Hilbert  
hilbysfe@gmail.com

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**Introduction:** Arterial brain vessel assessment is crucial for the diagnostic process in patients with cerebrovascular disease. Non-invasive neuroimaging techniques, such as time-of-flight (TOF) magnetic resonance angiography (MRA) imaging are applied in the clinical routine to depict arteries. They are, however, only visually assessed. Fully automated vessel segmentation integrated into the clinical routine could facilitate the time-critical diagnosis of vessel abnormalities and might facilitate the identification of valuable biomarkers for cerebrovascular events. In the present work, we developed and validated a new deep learning model for vessel segmentation, coined BRAVE-NET, on a large aggregated dataset of patients with cerebrovascular diseases.

**Methods:** BRAVE-NET is a multiscale 3-D convolutional neural network (CNN) model developed on a dataset of 264 patients from three different studies enrolling patients with cerebrovascular diseases. A context path, dually capturing high- and low-resolution volumes, and deep supervision were implemented. The BRAVE-NET model was compared to a baseline Unet model and variants with only context paths and deep supervision, respectively. The models were developed and validated using high-quality manual labels as ground truth. Next to precision and recall, the performance was assessed quantitatively by Dice coefficient (DSC); average Hausdorff distance (AVD); 95-percentile Hausdorff distance (95HD); and via visual qualitative rating.

**Results:** The BRAVE-NET performance surpassed the other models for arterial brain vessel segmentation with a DSC = 0.931, AVD = 0.165, and 95HD = 29.153. The BRAVE-NET model was also the most resistant toward false labelings as revealed by the visual analysis. The performance improvement is primarily attributed to the integration

of the multiscale context path into the 3-D Unet and to a lesser extent to the deep supervision architectural component.

**Discussion:** We present a new state-of-the-art of arterial brain vessel segmentation tailored to cerebrovascular pathology. We provide an extensive experimental validation of the model using a large aggregated dataset encompassing a large variability of cerebrovascular disease and an external set of healthy volunteers. The framework provides the technological foundation for improving the clinical workflow and can serve as a biomarker extraction tool in cerebrovascular diseases.

**Keywords:** artificial intelligence (AI), segmentation (image processing), UNET, cerebrovascular disease (CVD), machine learning

## INTRODUCTION

Stroke is one of the leading causes of death worldwide and 15 million people each year suffer a stroke (WHO EMRO, 2019). Stroke is a cerebrovascular disease and as such is characterized by changes in the arterial vasculature of the brain, such as stenosis and occlusion of vessels. Information about the arterial vessel status proves crucial for the diagnostic process. In chronic cerebrovascular disease—which is often present prior to stroke—it can serve as a biomarker: It was shown that brain vessel status can predict the likelihood of further stroke events (Gutierrez et al., 2015). In the acute clinical setting, brain vessel status provides stroke physicians with pivotal information. For example, acute arterial vessel occlusions qualify patients for mechanical thrombectomy, the best reperfusion therapy of stroke at the moment (Turc et al., 2019). This explains the high clinical relevance to depict brain vessels in cerebrovascular disease.

Non-invasive neuroimaging techniques are used to depict brain vessels. One method is magnetic resonance (MR) time-of-flight (TOF) imaging. It is fast, has no ionizing radiation exposure and can depict the arterial vasculature in high detail. Clinical reading of vessel imaging is based on visual judgment alone, as there is a lack of fully automated vessel segmentation methods. Fully automated segmentation means in the clinical context that the segmentation results would be rapidly available on the scanner console without the need of extra post-processing. Extensive, time-consuming and non-standardized image post-processing is an important obstacle to widespread clinical application. Fully automated analysis of TOF images could facilitate the diagnosis of vessel abnormalities and allow the quantification of the cerebrovascular status, e.g., arterial vessel density or arterial vessel diameters, to make potential biomarkers for diagnostic stratification easily available (Dengler et al., 2016; Santos et al., 2016; Yoo et al., 2018; Dutra et al., 2019; Murray et al., 2019).

Convolutional neural networks (CNNs)—a type of artificial neural networks (ANN) tailored for image analysis—have become the method of choice for vessel segmentation including the brain vasculature and showed promising results in recent applications (Moccia et al., 2018; Tetteh et al., 2018; Livne et al., 2019a). However, these previous studies suffer from various impediments that still limit their usability in the clinical setting.

First, post-processing of the images, such as noise filtration and brain masking are commonly applied (Passat et al., 2005; Livne et al., 2019a). The necessity of image processing may not only lead to suboptimal generalization due to a lack of standardized processing methods but may disqualify the models for clinical use due to time constraints, as the same processing needs to be applied to each image before predictions can be run. Second, the studies are often performed on small datasets or even simulated data due to a lack of large amounts of high-quality labeled data. Third, a lack of data heterogeneity limits the generalizability of these models to new datasets. Lastly, the suggested frameworks are rarely developed in cases with vessel pathologies. These constraints constitute a severe challenge for the clinical application of the models, since it may impair their ability to accurately detect vessels and vessel abnormalities in patients with cerebrovascular pathologies.

Recognizing this challenge, this work presents a high performance fully automated framework for vessel segmentation in patients with cerebrovascular disease addressing all the above-mentioned limitations, coined BRAVE-NET (BRAInVessel-NETwork). BRAVE-NET is a multiscale 3-D CNN model designed to dually capture high- and low-resolution volumes thus enabling enhanced distinction of brain vessels from other structures, such as skull areas and arteries not feeding the brain, making masking and other post-processing steps obsolete. The model was developed and validated using high-quality labeled data derived from multiple datasets of patients ( $n = 264$ ) with cerebrovascular disease. This approach ensured the high performance and high generalization of BRAVE-NET.

## METHODS

### Data Accessibility

Due to data protection laws, the imaging data used in this study cannot be published at the current time point. Implementation of the proposed network, as well as the training, prediction, and evaluation framework can be found on Github at <https://github.com/prediction2020/brain-vessel-segmentation>.

## Patients

Retrospective data from the PEGASUS (Martin et al., 2015), 7UP (Ultrahigh-Field MRI in Human Ischemic Stroke—a 7 Tesla Study), and 1000Plus (Hotter et al., 2009) studies, with 264 patients in total, were analyzed in this study. From the PEGASUS and 7UP studies, 74 and 9 patients with chronic steno-occlusive disease were available for this study, respectively. From the 1000Plus study, we included 181 patients with acute stroke. All mentioned studies were carried out in accordance with the recommendations of the authorized institutional ethical review board of Charité-Universitätsmedizin Berlin (PEGASUS and 1000Plus) and the Berlin state ethics boards (7UP). All subjects gave written informed consent in accordance with the Declaration of Helsinki.

## Imaging Properties

Time-of-Flight (TOF) Magnetic Resonance Angiography (MRA) images were used to train all benchmarked models. TOF relies on the fact that, within an imaged volume, inflowing blood has high magnetization compared to stationary tissue that becomes magnetically saturated by multiple radiofrequency pulses. TOF-MRA is one of the most important methods for non-contrast neurovascular and peripheral MRA. In the PEGASUS and 1000Plus studies, TOF imaging was performed on a Magnetom Trio 3T whole-body system (Siemens Healthcare, Erlangen, Germany) using a 12-channel receive radiofrequency (RF) coil (Siemens Healthcare) tailored for head imaging. In the 7UP study, TOF imaging was performed on a Magnetom Verio 3T whole body system (also Siemens Healthcare) using a 12-channel receive radiofrequency (RF) coil (Siemens Healthcare) tailored for head imaging. Parameters of the TOF imaging of each dataset are shown in **Table 1**.

## Data Labeling

Intra-cerebral arteries as well as the major brain-supplying arteries, i.e., the internal carotid arteries, the vertebral arteries and the basilar artery were labeled. For PEGASUS and 7UP data, ground-truth labels of the arterial brain vessels were generated semi-manually using a standardized pipeline. Pre-labeling of the vessels was performed by a thresholded region-growing algorithm using the *regiongrowingmacro* module implemented in MeVisLab (Mevis Medical Solutions, Bremen, Germany). To tackle inter-rater variability in label generation, these pre-labeled data were thoroughly manually corrected by either OUA and EA (both junior raters) and then cross-checked by the other rater. Junior raters were trained for image segmentation and were only allowed to independently label images once their performance met ground truth standards. These labels were then checked subsequently by VIM (9 years experience in stroke imaging). Thus, each ground-truth was eventually checked by three independent raters, including 1 senior rater. The total labeling time with this framework amounted to 60–80 min per patient. For 1000Plus data, the MeVisLab-based pre-segmentation step was replaced by using a 2D Unet segmentation model, developed in earlier work (Livne et al., 2019a). These pre-segmentations were—as described above—manually corrected either by OUA, EA, or JB, cross-checked by another junior rater

and then subsequently checked by VIM. For this approach, the total labeling time was reduced to 40–60 min per patient.

## Data Pre-/Post-processing

No pre- or post-processing methods were applied in the presented framework that are necessary to be performed on datasets prior to any prediction. Importantly, no mask was used for validation or test. All the developed models can operate on raw TOF images coming directly from the scanner and are not dependent on any processing step or external tool.

For the training step only, and here to reduce the training time of the models, a heuristical mask was automatically computed for each TOF image to exclude training patches containing only air as follows: (1) The images were smoothed with a  $16 \times 16 \times 16$  averaging filter in a shifting window fashion. (2) A threshold of intensity values at 10 was applied (intensity > 10 implies ROI). This provides an automatically generated mask for the training phase which contains all the brain tissue but also a thick border area around the skull. The inclusion of patches from these border areas proved to be crucial for the learning process, particularly for the distinction of the skull and the ability to capture the distinctive neighborhood of small vessels in that area. We note that this step serves the purpose of pre-selection of training patches only and is fully optional for the training phase. Inference with any of the models is possible on any raw image without a selection mask, consequently no masks were utilized in the testing phase. Similarly, instead of prior normalization of input data, an initial Batch-normalization layer was introduced to each architecture before other layers to learn and adjust to moving mean and variance statistics of training samples. This approach eliminates data pre-processing and the need of stashing statistics as well as enables quick adaptation when necessary through “fine-tuning” of Batch-normalization layers.

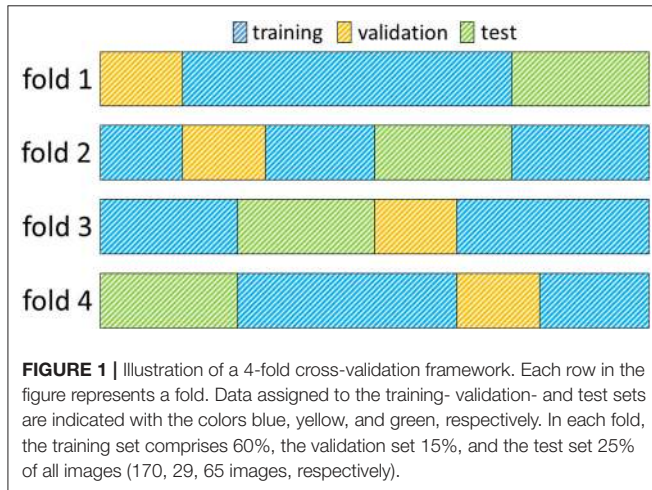
## Dataset Preparation and Patch Extraction

Images from each dataset were used for training, model selection and evaluation. A 4-fold cross-validation methodology was employed to ensure the robustness of all models toward different training and test sets. Therefore, data was split randomly into 4 distinct training and test sets making sure that each set contained the same number of patients from each source dataset (Pegasus, 7UP, 1000Plus). Furthermore, 15% of images were separated randomly without overlaps from each training set for validation and used for parameter selection. **Figure 1** illustrates the creation of training, validation and test sets. Final number of patients was 170, 29, 65 in each set, respectively; including equal numbers of patients from each source across folds.

The task of brain vessel segmentation on 3D MRI images challenges deep learning techniques from several aspects. First, processing of whole brain volumes at once requires significant resources in terms of GPU memory. Second, the distribution of vessels compared to brain tissue is very sparse. The physiological arterial vascular volume fraction of the brain is 1.5%. TOF-detectable arterial vessel voxels can be as low as on average 0.3% of all voxels within the brain. Finally, sufficient training of deep neural networks requires numerous images, while our dataset is—like most medical imaging datasets—limited in size.

**TABLE 1** | Time of flight (TOF) magnetic resonance imaging (MRI) parameters.

Dataset	Voxel size	Matrix size	TR/TE (ms)	Time of acquisition (min:s)	Flip angle (°)
PEGASUS	0.52 × 0.52 × 0.65	312 × 384 × 127	22/3.86	3:50	18
7UP	0.28 × 0.28 × 0.6	644 × 768 × 136	24/3.60	5:54	18
1000Plus	0.52 × 0.52 × 0.65	312 × 384 × 127	22/3.86	3:50	18



To solve these challenges, we redesigned the segmentation task as voxel-wise classification and trained our models on patches that represent an arbitrary neighborhood of the center voxel. Accordingly, 2,000 locations per TOF image (i.e., patient) were randomly sampled within the heuristical mask, ensuring 50% of the samples were vessel-centric. Patches of sizes  $[64 \times 64 \times 8]$  and  $[128 \times 128 \times 16]$  were extracted around each location, and comprised the input patches of our proposed framework.

### External Validation

In recent years, the medical AI research field showed a tendency to develop an overwhelming focus for performance in homogeneous samples in contrast to the requirements for a medical imaging product (Higgins and Madai, 2020). In the real world clinical setting algorithms are confronted with highly heterogeneous data reflecting different settings, multiple hardware options and various population properties. To this end, next to our—already heterogeneous—in-house dataset for model development and evaluation, we include an external validation set. We used the publicly available MRA TOF dataset of healthy volunteers from the MIDAS data collection website<sup>1</sup>, similarly to Bullitt et al. (2005). The dataset comprises images of 109 volunteers, evenly distributed into five age categories. We have chosen four images randomly from each age group, i.e., 20 in total—to thoroughly test generalization and robustness of our models but limit time requirements of the labeling process. Matrix sizes and voxel sizes were  $448 \times 448 \times 128$  and  $0.51 \times$

$0.51 \times 0.8$ , respectively. Other properties of image can be found on the original website. The same labeling procedure was applied as for 1000Plus data, described in the Data labeling section. Ground truth labels for the external validation are published under (Hilbert et al., 2020).

## Segmentation Frameworks

### Baseline Unet

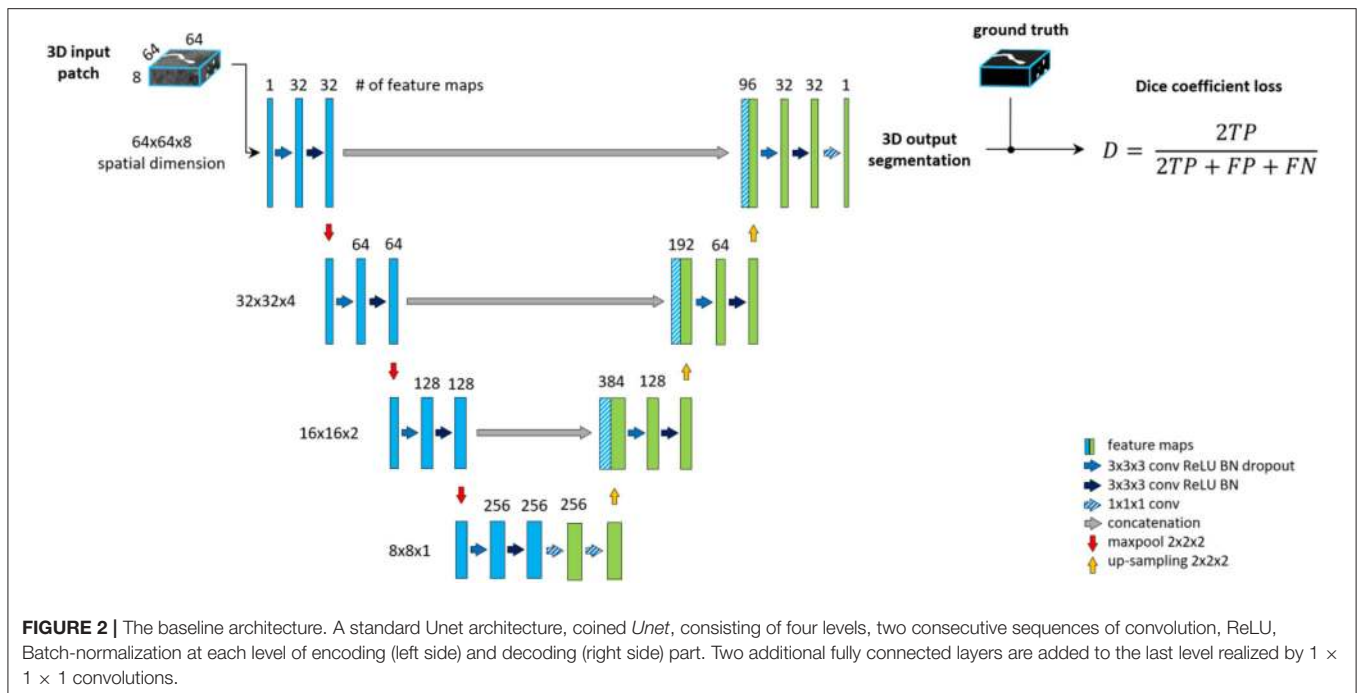
The backbone of the proposed framework is realized by an adjusted Unet architecture (Ronneberger et al., 2015) with four levels, shown on **Figure 2**. The Unet model consists of an encoding path (left side), and a decoding path (right side). At each level of the encoding, we employ a sequence of Convolution, Rectified Linear Unit (ReLU) activation and Batch-normalization two times consecutively followed by Max-pooling (Ioffe and Szegedy, 2015). Additional dropout is applied in the first sequence after the Batch-normalization (Srivastava et al., 2014). We set the kernels and stride to  $3 \times 3 \times 3$  and 1 for convolutions and  $2 \times 2 \times 2$  and 2 for max-pooling. The resolution and depth of feature maps are reduced to half and doubled, respectively after each level. As an extension of the original Unet architecture, we add two consecutive fully connected layers to the last level realized by  $1 \times 1 \times 1$  convolutions.

The decoding part of the architecture recovers the original input dimensions by applying the same sequences of Convolution, ReLU, Batch-normalization but replacing max-pooling with up-sampling at each level. Additionally, corresponding feature maps from the encoding part are concatenated to the input of each decoding level. Finally,  $1 \times 1 \times 1$  convolution with sigmoid activation is applied to map feature maps into a binary prediction map, which is used to calculate DSC loss.

### Context Unet

Patch-wise formulation of the segmentation problem comes at the cost of neglecting the full spatial context of input images. In theory, the model should learn to distinguish vessels from other structures within the brain. On the other hand, as described under the data labeling section, we are only interested in segmenting specific arterial brain vessels which have specific spatial locations. Furthermore, Livne et al. (2019a) showed the limitation of a standard Unet variant in detecting small vessels. A significant amount of these small vessels lie close to the skull, i.e., has a particularly distinctive spatial context. To tackle these problems, a multiscale approach is applied (Havaei et al., 2016; Kamnitsas et al., 2017; Yue et al., 2019). We extend the base architecture with a context aggregation under the hypothesis that it will improve the discrimination between vessels of interest and

<sup>1</sup><https://www.insight-journal.org/midas/community/view/21>



other constructs and particularly improve the segmentation of small vessels.

This approach is depicted in **Figure 3**. The encoding part of our Unet is extended with a so-called context path. The input of this path is a larger—context—patch, extracted around the same center voxel as for the other encoding path. Inspired by Kamnitsas et al. (2017), the input patch is then down-sampled by average-pooling with  $2 \times 2 \times 2$  kernels and stride of 2, i.e., to the same dimension but half-resolution compared to the other encoding path. The down-sampling allows for neglecting fine details and focusing on contextual information. The down-sampled input is fed into a parallel, equivalent sequence of layers. The two parallel downward paths are realized as duplicates (i.e., with no shared parameters) in order to enable distinctive feature encodings for the context and original patch. The output of the encoding paths, i.e., bottom level—are concatenated and fed through two fully connected layers realized by  $1 \times 1 \times 1$  convolution followed by ReLU. Finally, the residuals of each level of both encoding paths are concatenated to the input of the corresponding decoding level to facilitate the contribution of spatial and context information in the final prediction map.

### Deep Supervision Unet

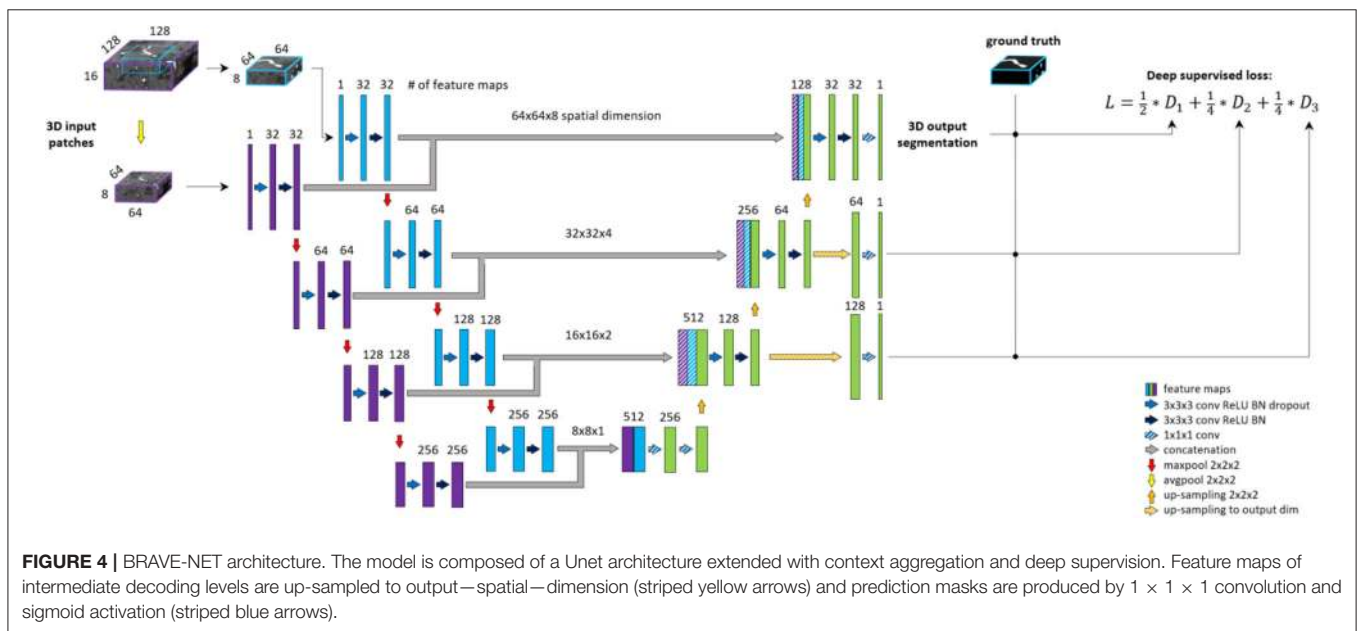
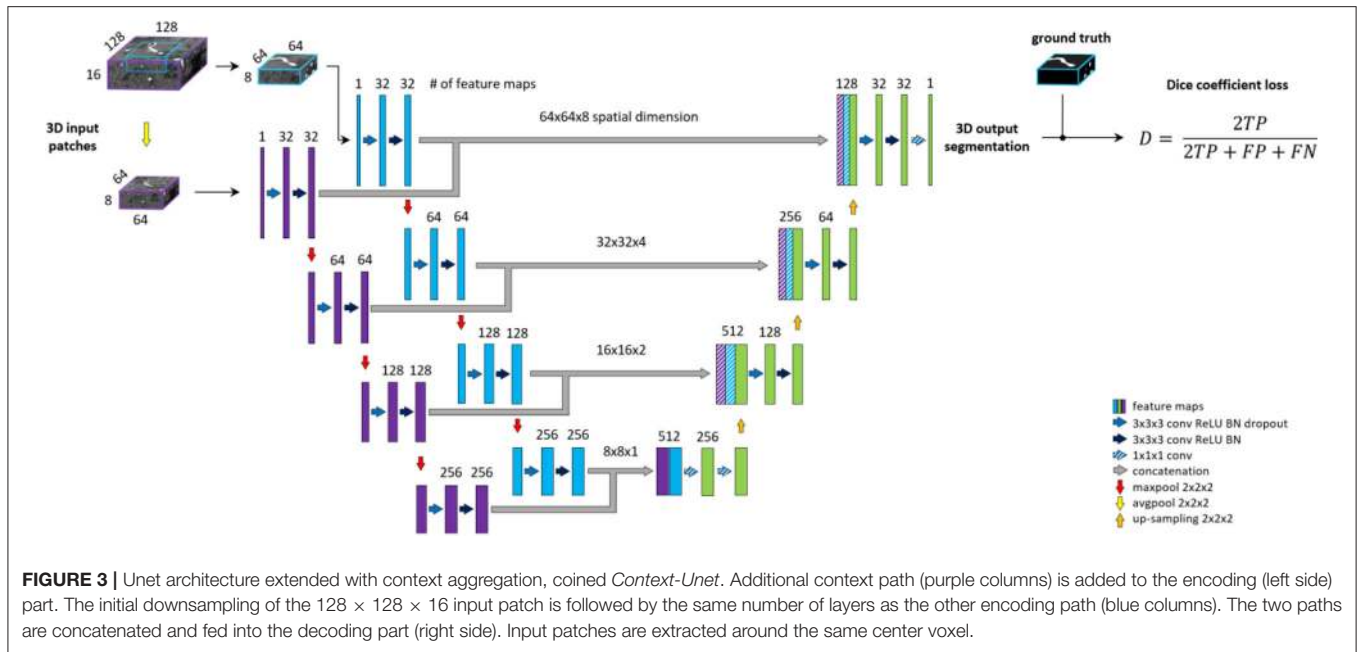
Deep supervision is a method commonly used to avoid the problem of exploding or vanishing gradients in deep networks by forcing intermediate layers to produce more discriminative features. Stawiaski (2017) implemented deep supervision by down-sampling ground truth segmentations and weighting each coefficient equally in the loss function. Other implementations aggregated the coarse, low-resolution feature maps into the final convolutional layers and thus incorporated different features in a single loss coefficient (Chen et al., 2014; Long et al., 2014; Folle

et al., 2019). In contrast, we aim to facilitate the convergence of intermediate layers by direct supervision. Feature maps from intermediate decoding levels (i.e., all except bottom and final level) are first up-sampled to the output dimension and then fed into a  $1 \times 1 \times 1$  convolutional layer with sigmoid activation to produce prediction masks. From each output of the model, DSC loss is computed with respect to the ground truth labeling. The loss coefficients are weighted and summed to create the training loss of the framework. We aim to reflect more emphasis on the final prediction thus assign 0.5 weight to the final layer and distribute the remaining 0.5 across intermediate outputs. A deep supervision variant of the architecture is coined here as DS-Unet. An illustration of deep supervision integration in the final architecture can be found in **Figure 4**.

### BRAVE-NET

The proposed BRAVE-NET architecture integrates the 3D Unet with both approaches introduced above, multiscale and deep supervision. **Figure 4** shows the BRAVE-NET architecture—including an illustration of the integration of deep supervision to the loss function. A model card outlining the model characteristics of BRAVE-NET can be found in **Appendix I**—Model card section of the Supplementary material, as suggested by Mitchell et al. (2019).

The number of trainable parameters are shown in **Table 2**. Naturally, Context-Unet and BRAVE-NET contain more parameters due to the multi-scale encoder part, while DS-Unet has only few more trainable parameters than Unet due to the two additional final convolutions of the up-sampled outputs. Mathematical formulations of encoder outputs and decoder inputs can be found in **Appendix III**—Mathematical formulations section of the Supplementary material.



## Model Training and Evaluation

### Experimental Setup

The models were trained in the above-described cross-validation framework. The optimal learning rate of the optimization process was selected by highest average performance across validation sets evaluated on whole brain segmentation. The largest batch-size feasible was used to fully utilize computational resources and to optimize training time. Models were trained on a high performance deep learning server using a single NVIDIA Titan RTX GPU.

To analyze the contribution of the proposed extensions, different configurations of the segmentation framework were tested. The performance of BRAVE-NET was compared to the DS-Unet, the Context-Unet, and to the baseline Unet.

### Training Scheme

The designated input (BRAVE-NET: [64 × 64 × 8], [128 × 128 × 16], DS-Unet: [64 × 64 × 8], Context-Unet: [64 × 64 × 8], [128 × 128 × 16], Unet: [64 × 64 × 8]) and ground truth patches ([64 × 64 × 8]) were used to train the models.

**TABLE 2** | Number of trainable parameters of all implemented models.

	Unet	DS-Unet	Context-Unet	BRAVE-NET
Number of trainable parameters	~6 million	~6 million	~10 million	~10 million

All models were implemented in Python using Tensorflow Deep learning framework (Abadi et al., 2015) and were trained using Adam optimizer (Kingma and Ba, 2014). The final layer of each model utilizes a voxel-wise sigmoid activation function, defined as  $p(x) = 1/(1 + \exp(f(x)))$ , where  $f(x)$  denotes the output value of the last  $[1 \times 1 \times 1]$  convolutional layer at the voxel position  $x \in \Omega$  with  $\Omega \in Z^3$  and  $p(x)$  is the approximated probability of a voxel  $x$  to be a vessel. The same activation is applied to each output of the deep supervision method. DSC loss was used to update the model parameters and to monitor training convergence, throughout all experiments. DSC between two binary volumes, i.e., prediction and ground truth—is defined as  $DSC = 2TP/(2TP + FP + FN)$ , where  $TP$  is the number of true positive voxels,  $FP$  is the number of false-positive voxels and  $FN$  is the number of false-negative voxels. Further derived:

$$\begin{aligned}
 DSC &= \frac{2TP}{(2TP + FP + FN)} \\
 &= \frac{2 \sum_{x \in \Omega} p_x g_x}{2(\sum_{x \in \Omega} p_x g_x) + (\sum_{x \in \Omega} p_x^2 - p_x g_x) + (\sum_{x \in \Omega} g_x^2 - p_x g_x)} \\
 &= \frac{2 \sum_{x \in \Omega} p_x g_x}{\sum_{x \in \Omega} p_x^2 + \sum_{x \in \Omega} g_x^2}
 \end{aligned}$$

Where  $p_x \in P: \Omega \rightarrow \{0, 1\}$  is the binary prediction volume and  $g_x \in G: \Omega \rightarrow \{0, 1\}$  the binary ground truth volume. This yields in the following differentiated gradient at each position  $j$ :

$$\frac{\partial D}{\partial p_j} = 2 \left[ \frac{g_j (\sum_{x \in \Omega} p_x^2 + \sum_{x \in \Omega} g_x^2) - 2p_j (\sum_{x \in \Omega} p_x g_x)}{(\sum_{x \in \Omega} p_x^2 + \sum_{x \in \Omega} g_x^2)} \right]$$

We monitored—patch-level—validation loss during training and employed EarlyStopping with a patience of 3 epochs to optimize training time and to save the best model with respect to validation loss. The weight of every convolutional layer was initialized with the commonly used Glorot uniform initialization scheme (Glorot and Bengio, 2010).

**Performance Evaluation**

The performance of vessel segmentation models can be assessed from many different angles. As Moccia et al. (2018) also highlights, different anatomical regions or imaging modalities might require different measures to be considered. To give a broader view of the performance of our models, we report five different metrics: Precision, Recall, DSC, 95 percentile Hausdorff Distance (95HD) and Average Hausdorff Distance (AVD). We included Precision and Recall to compare models by the quality and completeness of segmentations, respectively

and DSC, 95HD, and AVD to assess the spatial overlap between predictions and ground truth. The evaluation metrics were calculated on whole-brain segmentations (reconstructed from patch-wise predictions), by using the open-source evaluation tool from Taha and Hanbury (2015). We report mean values and standard deviation across test sets defined by the cross-validation framework. In case of the external validation set, we first generate ensemble predictions from each of the 4 models per architecture resulting from the cross-validation by averaging softmax scores voxel-wise then thresholding—using the same threshold of 0.5 as previously—into binary segmentations. Performance metrics are computed on these ensemble predictions.

**Visual Assessment**

For qualitative assessment, the predicted segmentation masks were transformed by in-house python code where true positives (TP), false positives (FP), and false negatives (FN) were assigned different voxel values (True negatives (TN) remained labeled with 0). The images were then visualized by overlaying these new masks with the original scans using ITK-Snap software (Yushkevich et al., 2006). By adjusting the opaqueness, it was possible to qualitatively assess which structures were correctly identified and which anatomical structures dominated with errors. Owing to the time burden of reviewing such a large number of images (4 models, 264 patients, 4-folds), only 1-fold was visually assessed. The first fold was chosen without prior inspection of any results. The 65 patients in the fold were assessed based on a predefined scheme slightly adapted from Livne et al. (2019a). Large vessels were defined as all parts of the internal carotid artery (ICA), the basilar artery, the anterior cerebral artery (ACA) and the M1, and P1 segments of the middle and posterior cerebral artery. All other parts were considered small vessels. The results of the visual analysis are summarized qualitatively in the results section. The scheme was the following: Vessel pathology detected (yes/no); Large vessels, overall impression (bad, sufficient, good); Small vessels, overall impression (bad, sufficient, good); Large vessels, errors (FP or FN dominating; or balanced); Small vessels, errors (FP or FN dominating, or balanced); Other tissue type segmentation errors (yes/no).

**RESULTS**

**In-house Dataset**

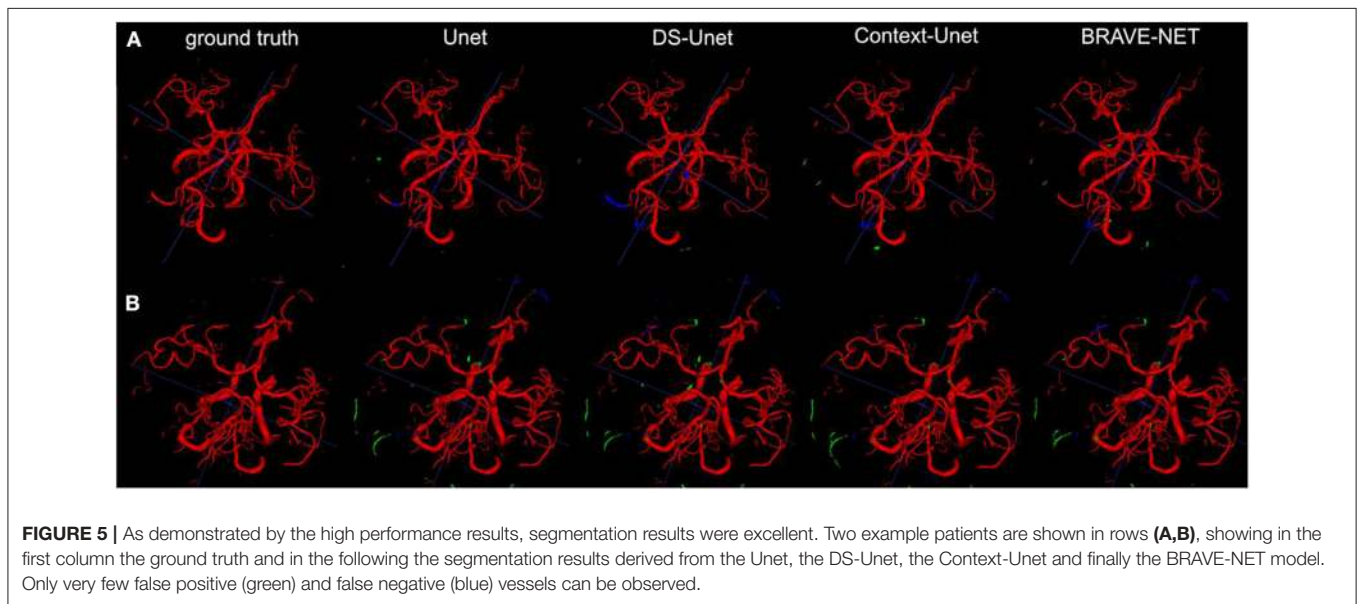
The BRAVE-NET model yielded the highest performance on all measures except Recall, namely DSC of 0.931, AVD of 0.165, 95HD of 29.153, and Precision of 0.941. Highest Recall of 0.930 was achieved by the Context-Unet model. The standard Unet baseline achieved a DSC of 0.928, AVD of 0.232, 95HD of 33.259, Precision and Recall of 0.928 and 0.929, respectively. Averaged test results of the cross-validation framework are presented in **Table 3**, while detailed results for each cross-validation fold can be found in Table 1 in **Appendix II**—Detailed results section of the Supplementary material. Example segmentation results can be found in **Figure 5**.

Context-aggregation alone was shown to improve the baseline performance of Unet for each measure while deep supervision

**TABLE 3** | Test results of cross-validation for all models, average (SD) over all folds are given. Best overall performance was achieved by our proposed BRAVE-NET model.

Models (all 3D)	DSC (SD)	AVD (SD)	95HD (SD)	Precision (SD)	Recall (SD)
Unet	0.928 (0.004)*	0.232 (0.041)*	33.259 (1.060)*	0.928 (0.005)*	0.929 (0.004)
DS-Unet	0.927 (0.004)*	0.222 (0.033)*	33.420 (0.866)*	0.927 (0.003)*	0.929 (0.005)
Context-Unet	0.931 (0.004)	0.198 (0.067)	29.279 (1.900)	0.934 (0.006)*	<b>0.930 (0.008)</b>
BRAVE-NET	<b>0.931 (0.003)</b>	<b>0.165 (0.013)</b>	<b>29.153 (0.988)</b>	<b>0.941 (0.005)</b>	0.923 (0.004)

An asterisk (\*) indicates significantly better performance of BRAVE-NET using the Wilcoxon Signed-Rank Test ( $z_{score} > z_{critical}$  at 5% significance level).



**FIGURE 5** | As demonstrated by the high performance results, segmentation results were excellent. Two example patients are shown in rows (A,B), showing in the first column the ground truth and in the following the segmentation results derived from the Unet, the DS-Unet, the Context-Unet and finally the BRAVE-NET model. Only very few false positive (green) and false negative (blue) vessels can be observed.

alone only improved the AVD measure. Combining both techniques however yielded the best overall performance.

For all models, the visual analysis found comparable excellent performance. Of the 65 patients, 15 showed visible stenocclusive changes. These vessel pathologies were detected by all models. The overwhelming majority of images was rated to have good performance both for large vessels (Unet: 95%; DS-Unet: 93%; Context-Unet: 95%; BRAVE-NET: 93%) as well as small vessels (Unet: 91%; DS-Unet: 91%; Context-Unet: 92%; BRAVE-NET: 92%). Large vessel error was dominated by false positives (~65% of images for all models), whereas small vessel error was dominated by balanced error (~63% of images for all models).

Besides excellent vessel segmentation performance, the visual inspection revealed a high rate of falsely labeled rare pathologies and other tissue classes. The false-labeling-rate was lowest for the BRAVE-NET model (40%), followed by Context-Unet (43%), DS-Unet (49%), and Unet (50%) with the highest false-labeling-rate. **Figure 6** demonstrated the improved performance of the BRAVE-NET model. Of all found false labelings, the majority were partially labeled segments of the external carotid artery in the neck region (48%), followed by false positively labeled deep veins (29%), as illustrated in **Figure 7**. Other false labelings were rarer [false labeling of hyperintensities in the mesencephalon 10%; cortical laminar necrosis 6%, cerebellar structures 2%,

arteria meningea media 2%, proteinaceous fluid (2%), and false negative labeling of rete mirabile structures 2%].

## External Dataset

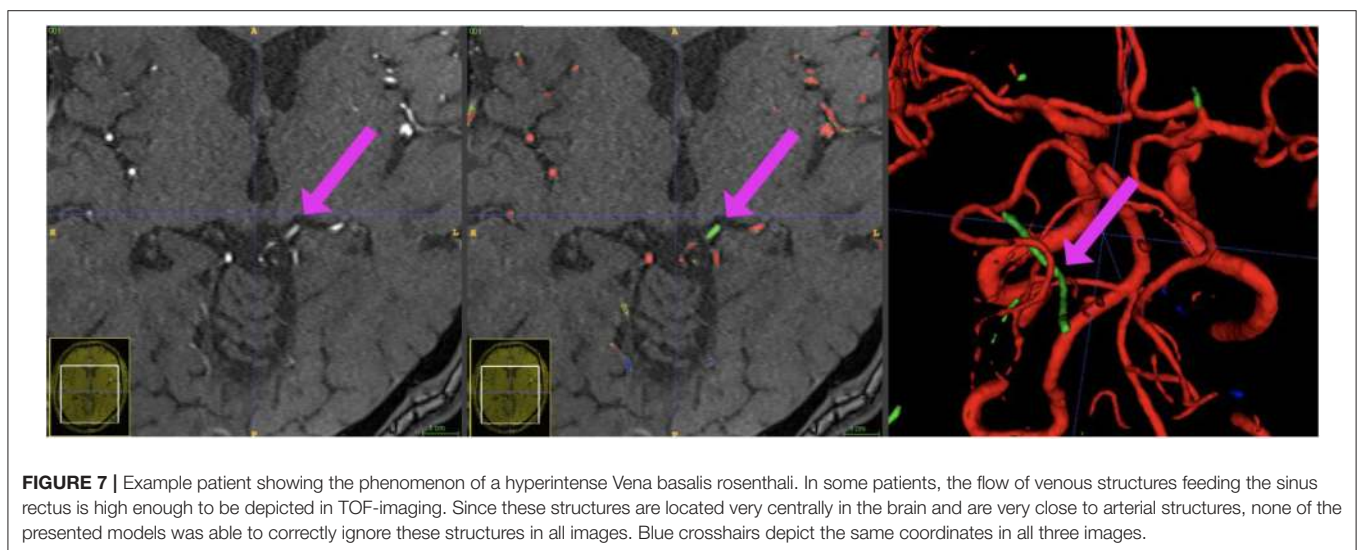
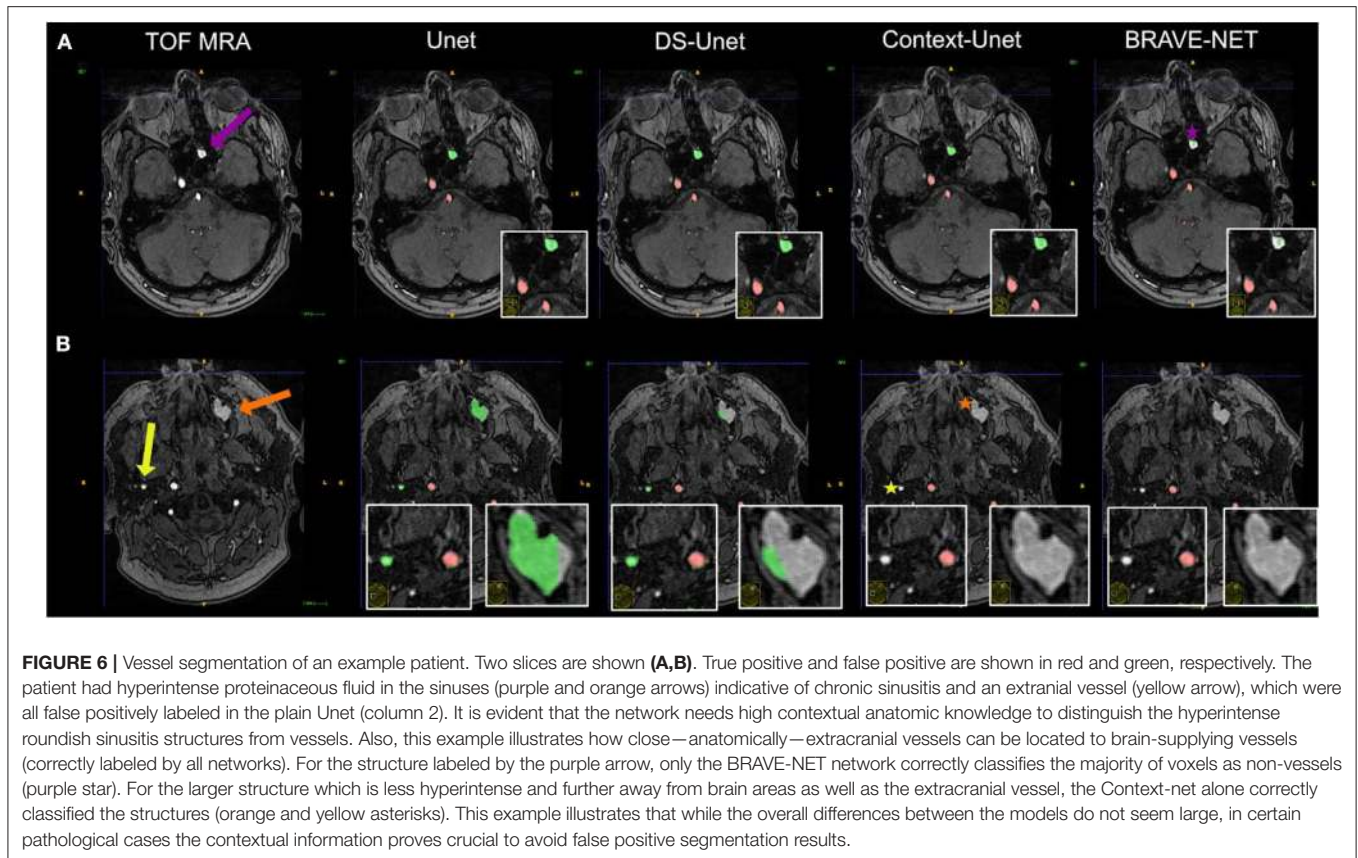
The results were consistent in case of our external validation set. BRAVE-NET achieved highest performance on all measures but Recall, where Context-Unet was superior. Details can be found in **Table 4**.

Visual inspection of the results revealed similarly high performance in terms of granularity and completeness of segmentations. The lower overall performance originated from the inclusion of large sinuses in the segmentations yielding a high number of false positives. As Precision values also indicate, the BRAVE-NET architecture was superior in reduction of false positives compared to other architectures. **Figure 8** demonstrates this in an example.

## DISCUSSION

We present the BRAVE-NET model for fully automated arterial vessel segmentation from TOF-MRA images in cerebrovascular disease. The model combines multiscaling and deep-supervision as extensions of the standard Unet approach and yielded a new state-of-the-art performance. This was achieved by robust





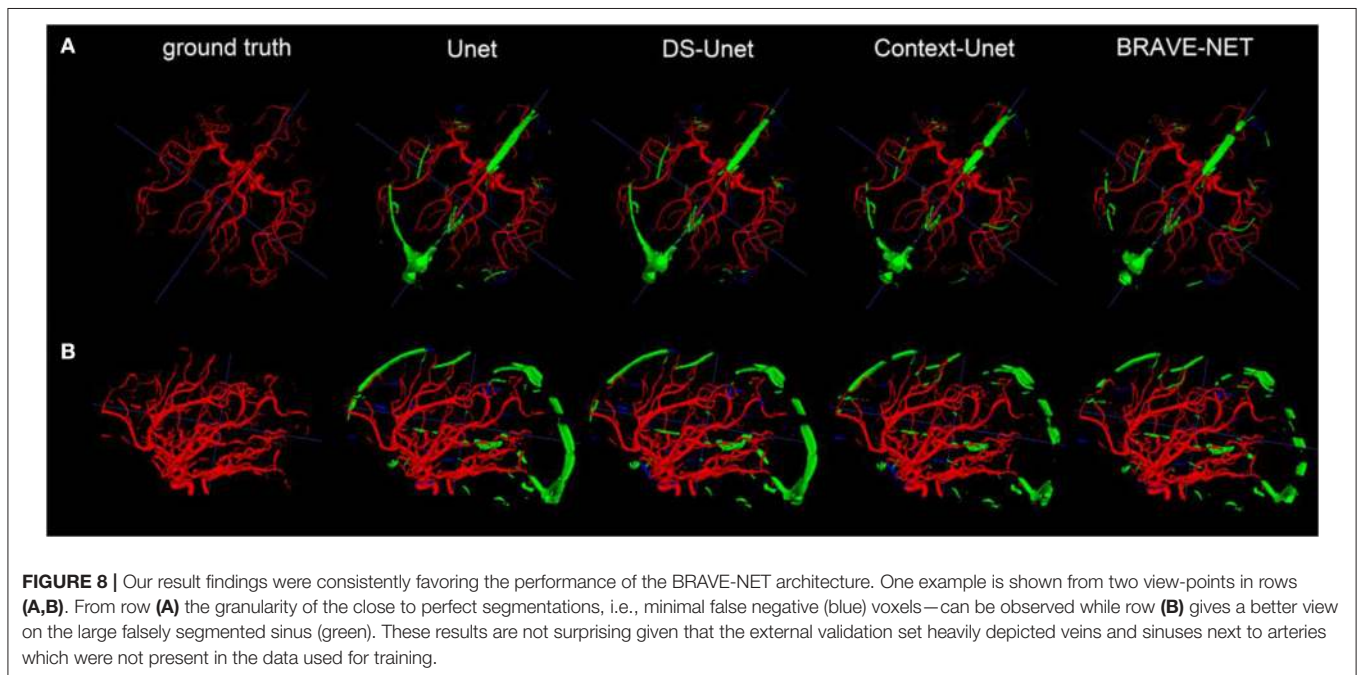
validation using the largest presented dataset in the literature so far. BRAVE-NET is suitable for clinical application, as it can be applied on raw images without the necessity for additional image processing. The superiority of the BRAVE-NET was demonstrated quantitatively and was confirmed via visual assessment. It outperformed the other tested models especially in cases of visible pathologies and false labeling. The performance

was further validated on an external set of healthy volunteers, where BRAVE-NET proved superior in quantitative measures as well as in exclusion of non-relevant brain structures (i.e., false positive reduction). BRAVE-NET marks a significant step toward increased applicability of deep learning based segmentations in the clinical setting and the continued development of quantitative vascular biomarkers in cerebrovascular disease.

**TABLE 4** | Test results on external validation set for all models, average (SD) performance across the 20 patients are shown.

Models (all 3D)	DSC (SD)	AVD (SD)	95HD (SD)	Precision (SD)	Recall (SD)
Unet	0.710 (0.038)*	2.044 (0.473)*	30.636 (7.157)*	0.659 (0.049)*	0.773 (0.044)
DS-Unet	0.719 (0.036)*	2.001 (0.443)*	29.647 (7.338)	0.665 (0.053)*	0.785 (0.031)
Context-Unet	0.745 (0.029)	1.689 (0.442)*	30.517 (7.641)*	0.700 (0.042)*	<b>0.798 (0.04)</b>
BRAVE-NET	<b>0.746 (0.036)</b>	<b>1.587 (0.461)</b>	<b>29.233 (7.027)</b>	<b>0.720 (0.049)</b>	0.778 (0.055)

Best overall performance was achieved by our proposed BRAVE-NET model. An asterisk (\*) indicates significantly better performance of BRAVE-NET using the Wilcoxon Signed-Rank Test ( $Z_{score} > Z_{critical}$  at 5% significance level). Bold values imply superior performance with respect to the certain metric.



Reported models in the literature for brain vessel segmentation are usually limited to a DSC of 0.90 (Babin et al., 2013; Wang et al., 2015b; Chen et al., 2017; Livne et al., 2019a). Notable exceptions, besides our work, are the works of Ni et al. (2020) and Patel et al. (2020) who reported DSC values of 0.96 and 0.94, respectively. They were, however, obtained for different modalities, CT angiography and digital subtraction angiography, respectively. Moreover, the DSC measure alone is not sufficient to provide the full picture regarding the needs of clinical application. Here, our model has other major advantages that have to our knowledge not been achieved together in one single model so far.

First, the majority of MRA-based vessel segmentation frameworks presented so far in the literature require image pre-processing, such as downsampling, brain masking, intensity correction, image normalization, and various other methods before they can be applied (Lesage et al., 2009; Chen et al., 2017; Livne et al., 2019a; Ni et al., 2020; Taher et al., 2020). Applications in computed tomography angiography (CTA) do not differ in that regard and include for example deformable matching, atlas co-registration, candidate vessel selection and feature extraction (Passat et al., 2005; Meijs et al., 2017). In contrast, our framework

completely avoids time- and resource-consuming pre- and post-processing methods that rely on domain knowledge. Moreover, this means that the BRAVE-NET model is usable on the scanner console in real-time given that a full brain prediction for clinically used dimensions, such as in the 1kplus study—can be available within minutes on a standard CPU system (AMD Ryzen 7 1700X), in our case 2 min. As a comparison, predictions for the same images on a standard GPU (NVIDIA Titan Xp) system take 40 s. Second, previous frameworks were developed using smaller datasets ranging from 10 to 100 patient scans derived from a single scanner (Passat et al., 2005; Wang et al., 2015b; Chen et al., 2017; Livne et al., 2019a; Ni et al., 2020; Patel et al., 2020). Taher et al. (2020) used a patient cohort size ( $N = 270$ ) similar to our study.

Third, the majority of reported models are based on data obtained from healthy individuals and did not include patients with (vessel) pathologies. Exceptions are the work of Meijs et al. (2017) who developed CTA-based models on data of patients with a suspected stroke, the work of Patel et al. (2020) including patients with aneurysms and the work of Livne et al. (2019a) who developed MRA-based models on patients with cerebrovascular disease and additionally tested the generalization

of the model on datasets from different scanners. The present work extends the latter work by adding a large number of patients from an additional data source allowing development, validation and testing on a combined multi-scanner, high quality labeled dataset. Furthermore, the model was developed on data with a broad range of pathologies present in patients with acute and chronic cerebrovascular disease. Thus, not only a broad variety of occlusions and stenoses were present, but also important pathologies, such as cortical laminar necrosis, rete mirabile, and others. Such a property is of high importance in the field of deep learning, as many applications suffer from bias due to constrained and homogenous data selection (Ho et al., 2019; Hofmanninger et al., 2020). Consequently, such models usually perform worse on datasets differing from the dataset distribution used for training. For example, Taher et al. (2020) used the freely available model of Livne et al. (2019a,b) for vessel segmentation on their own data and noticed a reduced performance compared to the original publication. Here, the increased generalizability of the model presented in this work is supported by its ability to capture a considerably greater degree of data variation caused both by larger technical and pathological variance in the datasets. Fourth, vessel segmentation provides a unique challenge owing to the complex nature of the vascular structure. Arterial brain vessels comprise around 1.5% of the input volume and therefore lead to an extreme class imbalance in the dataset. To mitigate this effect, a DSC loss function was used to neglect true negatives and subsampling of the training patches was applied so that 50% of all patches are centered around vessels ensuring representative and fair sampling. However, to properly evaluate vessel segmentation models, the performance measures must also reflect the vessel properties. Given ground-truth segmentations, accuracy, precision, and recall are the most frequent measures but numerous other measures exist. DSC is the most widely used metric and is therefore essential for examining the results with respect to the literature. Nonetheless, distance measures, such as AVD or 95HD have a distinct advantage over DSC in the context of vessels. Both DSC and HD measures are based on voxel-to-voxel comparisons between the segmented vessels and the ground-truth labels, without considering the continuity properties of the brain vasculature. However, since HD penalizes with respect to distance from the labeling it results in an implicit preference for (approximated) continuous structures. Previous studies have suggested HD measures as more useful to assess the performance of vessel segmentations (Taha and Hanbury, 2015). Thus, we assessed model performance based on five different measures: Precision, recall, DSC, AVD, and 95HD. This selection of metrics gives a comprehensive and complete overview of model performance as well as enables the presented state-of-the-art to be benchmarked thoroughly, i.e., beyond DSC—against future research. Fifth, one of the major strengths of our study is the validation of the models in patients with pathological conditions. Many pathologies are usually represented only by a small number of voxels compared to the full number of vessel voxels per patient. Some pathologies are present only in a small number of patients and are rare even in large samples. Therefore, these anomalies cannot be reflected properly by numerical performance measures but must

be assessed visually. Thus, we performed a thorough visual analysis of the results by a stroke imaging expert. Our findings indicated that the BRAVE-NET architecture outperformed the other models and was less likely to misclassify anomalies, such as proteinaceous fluid or laminar cortical necrosis. Nevertheless, the ratio of misclassifications, albeit often of minor markedness, was still 40% of all images. This underlines the importance of qualitative visual assessments to examine segmentation models for clinical applications. When quantitative metrics begin to plateau and stop showing meaningful differences, visual assessment is necessary to make clinically relevant distinctions. We, therefore, advise future studies to validate all proposed models both quantitatively and qualitatively, especially for the development of clinical applications.

Sixth, we included an external validation set to demonstrate the applicability of our framework with regards to no need of pre-processing steps, and test the generalization of our models on a public, out-of-sample dataset. Here, we made the contribution to generate 20 high quality ground truth labels for voxel-wise segmentation and make them publicly available (Hilbert et al., 2020). We advise future works to use these as an external test set in order to create a public benchmark for brain vessel segmentation. Our findings were shown to be consistent on this dataset. BRAVE-NET achieved superior performance in all but one measure further validating the benefits of the proposed improvements over the original Unet model. We also observed, however, a considerably lower overall performance. This can be attributed to the fact that the images from the external validation set show a heavy depiction of venous structures (internal veins and especially brain sinuses) next to arterial vessels. As our training set contained almost exclusively arteries, it is no surprise that the models struggled with these out-of-sample structures. These findings highlight the heterogeneous nature of real world data and that the development of tools with very broad applicability in the clinical setting require a dedicated validation strategy encompassing as many variations of input data as possible (Higgins and Madai, 2020).

To summarize, the presented framework add to the literature through the unique combination of high performance, full automation and robust validation. It thus has the largest applicability for the clinical setting so far.

The excellent performance of BRAVE-NET can also be attributed to architecture design choices. CNNs and encoder-decoder Unet variations, in particular, were proven in recent years as the state-of-the-art method for semantic segmentation including medical imaging applications (Milletari et al., 2016; Badrinarayanan et al., 2017; Dong et al., 2017; Alom et al., 2018; Norman et al., 2018; Huang et al., 2020; Sinha and Jose, 2020; Zhou et al., 2020). This success was further demonstrated in vessel segmentation and specifically in brain vessel segmentation problems (Chen et al., 2017; Alom et al., 2018; Huang et al., 2018; Livne et al., 2019a; Yue et al., 2019). The utilization of the Unet as a baseline architecture was, therefore, a natural choice. Nonetheless, the classic Unet architecture was previously shown to be prone to errors in the presence of visible pathologies especially in segmenting small vessels and with regards to the skull (Livne et al., 2019a). To target these issues

we aimed to improve spatial information integration using multiscale- and 3-D variants. Multiscale architectures, i.e., fusion of information from multiple resolutions—were shown to be beneficial in boundary detection and segmentation problems (Kokkinos, 2015; Havaei et al., 2016; Kamnitsas et al., 2017; Stawiaski, 2017; He et al., 2019; Sinha and Jose, 2020). Some recent works proposed re-thinking of skip-connections within the Unet architecture to effectively match semantics and fuse information on multiple resolutions for prediction (Huang et al., 2020; Zhou et al., 2020). Others argue that multiscaling in encoder-decoder architectures lead to the use of redundant information and rather propose integration guided self-attention to extract relevant features from more discriminant regions (Sinha and Jose, 2020). We implemented multiscaling following a common practice across applications, namely employing an additional encoder path of the architecture operating on lower resolution. This so-called context path, allowed to capture the broader anatomical context of the vessels and particularly target the detection of small vessels. Multiscale integration was also important to tackle the challenge of unmasked images: it allowed distinguishing larger anatomical structures, such as the skull and neck areas from the regions with brain and brain-supplying vessels while maintaining high resolution of the segmented vessels. The down-sampling within the context path enabled to avoid redundant extraction of features while maintaining corresponding spatial integrity of the parallel paths. Multiscale models yielded improved performance in all measures compared to the baseline Unet, corroborating the advantage of multiscaling in semantic segmentation tasks. Moreover, deep supervision in CNNs was previously shown to be advantageous as a strong “regularization” for classification accuracy and learned features and tackling problematic convergence behavior (Lee et al., 2015). Different variations of deep CNNs were described in the literature and demonstrated improved results and faster convergence (Lee et al., 2015; Szegedy et al., 2015; Wang et al., 2015a; Dou et al., 2016). In our study, deep supervision improved performance only in combination with the multiscaling approach.

Our study has several recognized limitations. First, while the dataset in this study was acquired from multiple scanners and patients exhibited a large spectrum of steno-occlusive cerebrovascular disease, the data heterogeneity was still limited. It is highly warranted to specifically include in future work datasets with other pathologies, such as aneurysms, other MR scanner- and MR imaging sequence types and other imaging modalities, such as CT. Here, transfer learning approaches seem beneficial. Heterogeneity is not only essential in training sets but also in testing for thorough and robust assessment of performance. Even though our external validation set yielded consistent ranking of our models, segmentation performance was rather limited due to a high false positive rate throughout all architectures. From a development perspective, we expect this to be easily overcome by minor fine-tuning on a few samples from the dataset, and this highlights that all variations of input data need to be used for tools with planned clinical applications. Second, it was beyond the scope of this work to integrate all potentially beneficial architecture modifications. Further future modifications could utilize densely connected convolutional

blocks to enhance feature propagation through encoding levels (Zhang et al., 2018) or even a more complex Adaptive Pyramid Context Network with local affinity integration to advance multiscale context aggregation using global representation (He et al., 2019). Moreover, the Unet++ architecture shown in Zhou et al. (2020) would seem like a well fit candidate for direct comparison. Unet++ realizes multi-scale processing through nested, dense pathways between the encoder and decoder levels alleviating the fusion of semantically dissimilar features. The applied Deep Supervision on the dense, up-sampling pathways from intermittent layers of the encoder enables learning of lower resolution features important for prediction, similarly to our context-path. Third, due to well-recognized resource constraints only 1-fold was reviewed visually. It is a general challenge for this kind of research that quality characteristics, such as cross validation and large data sets lead to an increasing amount of images that require highly time-consuming visual review. It is unlikely that reviewing more folds would lead to major changes in the visual assessment, but it cannot be ruled out with certainty.

## CONCLUSION

We present a fully automated deep learning framework for TOF-MRA brain vessel segmentation. We utilize multiscaling for better spatial information integration and deep supervision for improving model convergence. We employ a multi-scanner dataset of yet unmatched size and range of pathologies in the published literature. We also ensure a high degree of robustness of our framework through extensive quantitative and qualitative evaluation. We explicitly analyze and communicate model performance in presence of relevant pathologies to further increase applicability in the clinical setting. Our results thus constitute a new state-of-the-art for fully automated frameworks for brain vessel segmentation. Our work strongly facilitates and promotes the usability of automated vessel segmentation in the clinical setting and can be swiftly translated to the development of cerebrovascular biomarkers for clinical applications.

## DATA AVAILABILITY STATEMENT

The datasets used in this article for model training are not readily available because data protection laws prohibit sharing the 1000Plus, PEGASUS and 7UP datasets at the current time point. Requests to access the datasets should be directed to [ethikkommission@charite.de](mailto:ethikkommission@charite.de). The imaging data used for external validation and the generated labels are available under <https://www.insight-journal.org/midas/community/view/21> and <http://doi.org/10.5281/zenodo.3968844> respectively.

## ETHICS STATEMENT

The studies involving human participants were reviewed and approved by Ethics committee of Charité University Medicine Berlin and Berlin State Ethics Board. The patients/participants provided their written informed consent to participate in this study.

## AUTHOR CONTRIBUTIONS

AH, VM, AT, JF, DF, and ML: conceptualization. JS, JW, PD, TN, JF, and DF: funding acquisition and project administration. EA, OA, JB, VM, AK, IG, and JF: data curation. AH, VM, JS, IG, AK, AT, JW, PD, TN, JF, DF, and ML: methodology. AH, VM, DF, and ML: model development and validation. JS, JW, TN, JF, and DF: resources. AH, VM, DF, and ML: writing—original draft. AH, VM, EA, OA, JB, JS, IG, AK, AT, JW, PD, TN, JF, DF, and ML: writing—review and editing. All authors contributed to the article and approved the submitted version.

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## SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: <https://www.frontiersin.org/articles/10.3389/frai.2020.552258/full#supplementary-material>

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**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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