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# Carotid atherosclerotic plaque stability prediction from transversal ultrasound images using deep learning

Predikce stability aterosklerotického plátu z transverzálních ultrazvukových obrazů pomocí hlubokého učení

#### **Abstract**

Aim: To automatically predict the stability of carotid artery plaque from standard B-mode transversal ultrasound images using deep learning. A reliable predictor would reduce the need for follow-up examination and pharmacological and surgical treatment. Methods: A region of interest containing the carotid artery was automatically localized. An adversarial segmentation method was trained on a combination of a small pixelwise annotated dataset and a larger weakly annotated dataset. A multicriterion regression with automatic weight adaptation was applied to predict a series of clinically relevant attributes, including the plaque width increase over 3 years. Results: The current plaque width could be estimated with a high correlation ( $\rho = 0.32$ ) and a very high statistical significance. The estimated future increase of the plaque width was correlated less ( $\rho = 0.22$ ) but statistically significantly (P < 0.01). The correlation between automatic and expert assessments of echogenicity, smoothness and calcification was even smaller. Conclusion: We confirmed a relationship between the plaque appearance in ultrasound and the probability of its future growth, but it is too weak to be used in clinical practice as the sole predictor of the plaque stability.

#### Souhrn

 $\it Cil:$  Automaticky předpovídat stabilitu aterosklerotického plátu v karotidě ze standardních transverzálních ultrazvukových obrazů v B-modu za použití hlubokého učení. Spolehlivý prediktor by snížil potřebu klinických kontrol i farmakologické či chirurgické léčby.  $\it Metody:$  Automaticky byla lokalizována oblast zájmu obsahující karotidu. Adversariální metoda segmentace byla natrénována na kombinaci malého kompletně anotovaného datasetu a většího slabě anotovaného datasetu. Multikriteriální regrese s automatickou adaptací vah byla použita k predikci série klinicky relevantních atributů, vč. nárůstu tloušťky plátu během 3 let.  $\it Výsledky:$  Současnou šíři plátu bylo možno odhadnout s vysokou korelací (p=0,32) a velmi vysokou statistickou signifikancí. Odhadovaný budoucí nárůst šíře plátu byl korelován méně (p=0,22), ale stále statisticky významně (p<0,01). Korelace mezi automatickým a expertním hodnocením echogenicity, hladkosti a kalcifikací byla ještě nižší.  $\it Závěr:$  Potvrdili jsme závislost mezi vzhledem plátu v ultrazvukovém obraze a pravděpodobností jeho budoucího růstu, ale je příliš slabá, než aby byla využitelná v klinické praxi jako jediný prediktor stability plátu.

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# Klíčová slova

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#### Introduction

Carotid artery atherosclerotic plaques are present in over half of the population at the age of 50 [1]. In most cases, the plaques are stable, asymptomatic, and present no health hazard. However, some plaques keep growing rapidly and may lead to thrombosis or thromboembolism with possible blockage of the blood supply to the brain.

Atherosclerotic plaques can be studied by histology [2], CT, MRI [3], Doppler ultrasound [4] or other methods. Nowadays, the vast majority of carotid artery atherosclerotic plaques are detected by standard B-mode ultrasonographic screening [5]. This work aims to use these B-mode images to predict whether the plaque is stable (and will not grow substantially) or unstable and needs to be treated more aggressively. A success would have a profound effect in reducing the need for follow-up examinations, pharmacological treatment, and surgery.

Previous studies have shown that atherosclerotic plaque stability is related to some manually or semi-automatically evaluated characteristics derived from the ultrasound image such as the plaque width, surface irregularity and exulceration [6,7], echogenicity and homogeneity [8], neovascularity and complexity [9], or texture features [10]. However, the association found was weak, not allowing to make diagnostic decisions. Moreover, there does not seem to be a consensus concerning relevant features. Some studies even found no significant relationship [11].

The goal of this work is to automatically predict the stability of carotid artery plaque

from standard B-mode transversal ultrasound images using deep learning. We have previously shown a fully automatic image analysis pipeline [12] calculating geometry and wavelet features which were significantly different (p  $< 10^{-3}$ ) between stable and unstable plaques, but the accuracy of the predictor remained low (61–62%). Here we replace classical image analysis algorithms by deep learning methods, using several new techniques for this task, and hope to outperform the previous methods.

First, we employ adversarial learning techniques to learn to segment also from the electronic caliper images instead of only from full pixel-wise annotations. Second, we use multicriterion regression with automatic weight adjustment to predict not only the plaque width increase but also other parameters determined by the sonographers. We combine the segmentation mask and the original image as inputs to a relatively shallow multi-channel regression network. All techniques help to avoid overfitting and reduce the required annotation effort.

# Materials and methods Dataset

We used data from the clinical study AN-TIQUE [13], from which we extracted images of 413 patients with a median age of 69 years who had an atherosclerotic plaque localized in the carotid bifurcation or proximal part of the internal carotid artery (ICA) with a width 2.0 mm or more and a degree of stenosis at least 30% in at least one exami-

nation. The patients were examined approximately every 6 months over the period of 3 years between 2015 and 2019 using Mindray DC8 ultrasound scanner (Mindray, Shenzhen, China) with a linear 3–12 MHz probe. All patients provided an informed consent and the images were anonymized before further processing. The images were categorized as transversal, longitudinal, conical and Doppler [12]. In this work, only transversal images were used. Two sets of images from two examination waves were analyzed for each patient – one immediately after enrollment and the second after 3 years.

#### **Image attributes**

The experts used one key image to determine the plaque width (denoted d) in millimeters using virtual calipers as well as a set of image attributes describing the plaque appearance, such as echogenicity or homogeneity (see Tab. 1 for a full list). Our main prediction target was the positive part of the plaque width increase between the two examination waves:

 $\Delta = \max (d_2 - d_1, 0).$ 

The full vector of the attributes for image j will be denoted  $v_j = (v_j^1, v_j^2, ..., v_j^A)$  with A = 9,  $v_j^1 = d$  being the plaque width and  $v_j^2 = \Delta$  the width increase, etc.

In this study, we only used key images to avoid discrepancies between the images and the expert annotations. For each patient and each side (left and right), we therefore had two images, called first and second. See Fig. 1 (left) for an example.

Tab. 1. Image attributes assigned by experts and the results of their automatic prediction — correlation coefficient  $\rho$ , P-value and the RMSE. Dashes indicate the prediction was not significantly correlated (P > 0.05).

			prediction			
symbol	name	units	range	ρ	P-value	RMSE
$v_j^1 = d$	plaque width	mm	0-8	0.32	< 10 <sup>-7</sup>	0.88
$V_j^2 = \Delta$	width increase	mm	0–2	0.22	0.008	0.52
$V_j^3$	surface smoothness		1–3	0.13	0.04	0.64
V <sub>j</sub> <sup>4</sup>	echogenicity		1–4	0.12	0.05	1.03
$V_j^5$	homogenicity		1–2	_	_	_
V <sub>j</sub> 6	low echo area	%	0-100	_	_	_
$V_j^7$	mid echo area	%	0-100	_	_	_
V <sub>j</sub> <sup>8</sup>	high echo area	%	0-100	_	_	_
V <sub>j</sub> <sup>9</sup>	calcification area	%	0–100	0.13	0.04	23.5
RMSE – root mean square error						

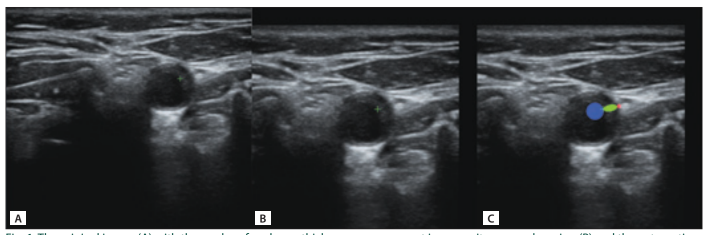


Fig. 1. The original image (A) with the markers for plaque thickness measurement in green, its cropped version (B) and the automatically created incomplete weak segmentation (C).

Obr. 1. Původní snímek (A) se značkami pro měření tlouštky plátu zeleně, jeho oříznutá verze (B) a automaticky vytvořená neúplná segmentace (C).

#### Localization

For each transversal image, the lumen of the ICA was automatically localized using a Faster R-CNN architecture [14] trained on 300 manually annotated images, as described in [12]. A region of interest (ROI) centered over the carotid artery of size  $520 \times 520$  pixels was extracted. See Fig. 1 (center) and 2 (left) for examples.

For each key image, an image with the distance markers (virtual calipers) in green was available (Fig. 1). The cross marker is located at the interface between the plaque and the lumen and a dashed line connects it with the vessel wall. We located the cross by template matching in the green channel and the wall interface point was determined as the furthest green point from the marker. Note that markers were not used for the ROI localization or any other processing, so our method is not restricted to key images.

From the 413  $\times$  2  $\times$  2 = 1,652 potential key images, 1,399 were available after ROI and marker localization described above. These were randomly split into training (80%) and testing (20%) part, yielding 568 training and 147 testing images to estimate  $\Delta$  (from the first examination wave only) and 1 113 training and 286 testing images to estimate the plaque width d and all other attributes (from both examination waves).

#### Segmentation

To train the segmentation network, we used two sets of annotations, strong and weak. In the smaller, strongly annotated dataset  $\Omega_s$ , experts assigned each pixel to one of m = 4 classes (1 – background, 2 – ves-

sel wall, 3 - plaque, and 4 - lumen) [12], see Fig. 2. There were 111 training (80%) and 27 testing images (20%), distinct from the key images described above. To increase the segmentation dataset size, we leveraged the already identified distance markers. We put an ellipsoidal mask with an aspect ratio 1:4 for the plaque class over the line connecting the two markers and circular masks with the radii 20 and 5 pixels touching the line from the outside for the lumen and wall classes, respectively (see Fig. 3, right). This way, we obtained weak segmentation annotations for all 1,399 key images without requiring any additional timeconsuming manual annotation. We denoted the dataset  $\Omega_{w}$ . The annotations are called weak because only a small part of the pixels is annotated, which can nevertheless guide the segmentation to the correct location of the plaque. Both strong and weak annotations can be represented by onehot encoding, using an indicator vector y<sub>i</sub> =  $(y_i^1, y_i^2, y_i^3, y_i^4) \in \{0,1\}^m$  for each pixel i, with  $y_i^1=1$  indicating background,  $y_i^2=1$  the vessel wall etc. For strong annotations the class is always determined,  $\sum_{k} y_{i}^{k} = 1$ , while for weak annotations in undetermined pixels  $y_1 = (0,0,0,0).$ 

The segmentation convolutional neural network (CNN) S uses a standard U-net [15] architecture with Resnet34 [16] backbone and softmax output layer with an m = 4 channel output f in the range [0,1]. The network was trained on a union  $\Omega = \Omega_{\rm s} \cup \Omega_{\rm w}$  of strongly and weekly annotated datasets, minimizing the cross-entropy segmentation loss function

$$L_{seg} = -\underset{j \in \Omega}{\text{mean mean}} \sum_{k=1}^{m} y_i^k logf_i^k (x_j)$$

where the indices j, i, k correspond to training images, pixels and classes, respectively. This way, the undetermined pixels in the weakly annotated images are automatically ignored. The strongly annotated dataset is oversampled 15x to give it more weight. To combat overfitting, we used a wide range of augmentation operations: random horizontal flipping, small translations, rotations and scaling changes, changes of brightness and contrast, and multiplicative and additive noise.

Adding the weakly annotated dataset improves the segmentation results with respect to using only the small strongly annotated dataset [12]. However, we found that the U-net nevertheless sometimes produced unrealistic and physiologically unfeasible segmentations. As a remedy, we used generative adversarial nets (GANs) [17] to learn the model of likely segmentations [18]. Besides the segmentation network S, we therefore also trained a discriminator D, which should learn to distinguish 'real' human expert segmentations from the strongly annotated dataset on one side, from 'fake' segmentations produced by S on both strongly and weakly annotated images on the other side. The segmentation network S then learns to generate plausible segmentations which are accepted by the discriminator D as "real".

We trained the segmentation network S to minimize the combined loss function

$$L_{S} = \alpha L_{seg} - \underset{j \in \Omega}{\text{mean logD(S(x_{j}))}}$$

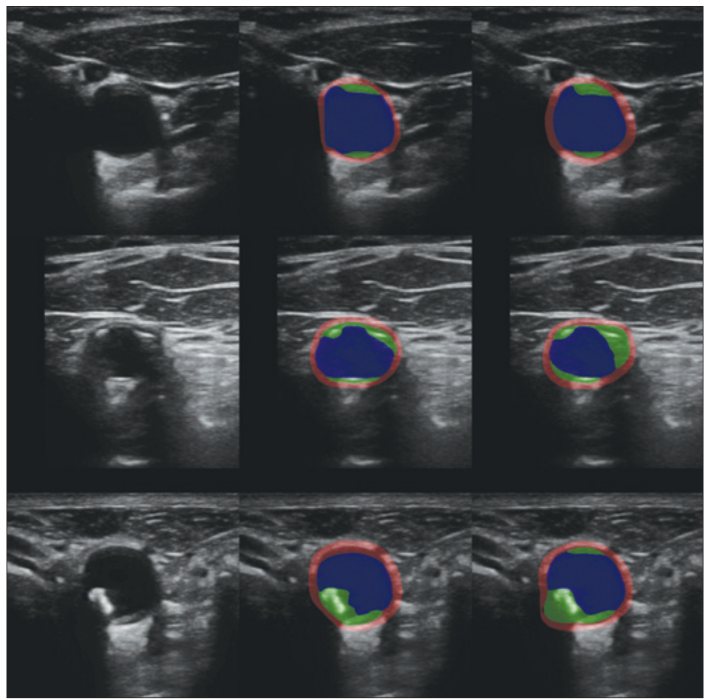


Fig. 2. Example segmentations on the test part of the strongly labeled dataset. From the left: input image, expert segmentation, automatic segmentation.

Obr. 2. Příklady segmentace na testovací části plně označeného datasetu. Zleva: vstupní obrázek, expertní segmentace, automatická segmentace.

which rewards S for generating segmentations that are evaluated by the discriminator D as "real", corresponding to D(S(x<sub>i</sub>))  $\approx$  1. The coefficient  $\alpha$  controls the trade-off between the segmentation error and the prior. We used  $\alpha=$  100, the exact value does not seem to be critical.

In contrast, the discriminator D was trained to minimize

$$L_{_{D}} = -1 \; / \; |\Omega| \left( \sum_{j \; \in \; \Omega_{_{S}}} log \; D(y_{_{j}}) + \sum_{j \; \in \; \Omega} log \; (1 \; - \; D(S(x_{_{j}}))) \right)$$

encouraging D to classify all expert segmentations from  $\Omega_{\rm s}$  as 'real' (D  $\approx$  1) and all segmentations generated by S (from both  $\Omega_{\rm s}$  and  $\Omega_{\rm w}$ ) as 'fake' (D  $\approx$  0).

It is notoriously difficult to prevent the discriminator D to work too well and thus de-

prive the segmentor from useful gradient feedback, especially in the beginning, when S is not well trained. To make the task more difficult for the discriminator, we used three strategies. First, we added label noise (randomly flipping the real/fake label with probability 0.1) and we also degraded the segmentations by adding a uniform noise with amplitude 0.3 to the class probabilities y<sub>i</sub> and

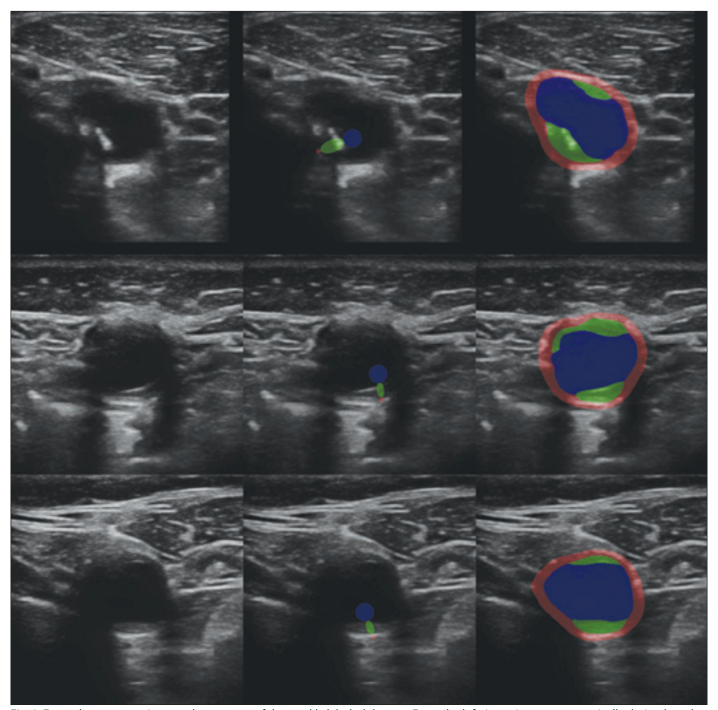


Fig. 3. Example segmentations on the test part of the weakly labeled dataset. From the left: input image, automatically derived weak segmentation, automatic dense segmentation.

Obr. 3. Příklady segmentací na testovací části datasetu s neúplným označením. Zleva: vstupní obrázek, automatická neúplná segmentace, automatická plná segmentace.

 $S(x_j)$  and blurring them with a Gaussian filter with  $\sigma=3$  "pixels" . The probabilities were renormalized to sum to unity afterwards. Second, we switched between optimizing S and D based on the performance of the discriminator D. When its accuracy in distinguishing 'real' from 'fake' segmentations exceeded 60%, we switched to optimizing the segmentor S. When the accuracy dropped

below 40%, we stoped optimizing S and started optimizing D. We used the Adam optimizer with a relatively small step size 10<sup>-5</sup>.

Third, the discriminator CNN D is very small. It is a simplified version of the discriminator network from [19], using 4 convolutional layers with  $3 \times 3$  kernels and 16, 32, 64 and 128 channels, respectively. We used leakyReLU non-linearities, dropout

(with probability 0.25), and instance normalization after every convolutional layer. The final linear layer with sigmoid nonlinearity produces the desired output value  $D(y) \in [0,1]$ .

Examples of automatic segmentations are shown in Fig. 2 and 3 (right column) for the strongly and weekly annotated images, respectively.

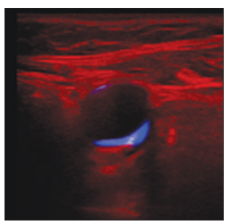


Fig. 4. An example of three channel input for the regression network shown as an RGB image: the input image in red, its masked plaque region in green and the soft mask itself in blue.

Obr. 4. Příklad tříkanálového vstupu pro regresní síť zobrazeného jako barevný obrázek: vstupní obrázek červeně, jeho maskovaná oblast plátu zeleně a samotná maska modře.

#### Regression

The image attributes v<sub>j</sub> to be predicted are linearly normalized to the range [0,1] (see Table 1 for the original ranges) for better numerical stability. We chose to minimize a weighted mean squared error (MSE):

$$L_{reg} = mean \sum_{j=1}^{A} W_{i} (\hat{V}_{j}^{i} - \tilde{V}_{j}^{i})^{2}$$

where  $\tilde{v}_j^i = (\tilde{v}_j^1 \dots \tilde{v}_j^A)$  are the normalized expert-determined attributes,  $\hat{v}_j^i = (\hat{v}_j^1 \dots \hat{v}_j^A)$  = R(u<sub>j</sub>) are the normalized predictions and R is the prediction network with input u<sub>j</sub> associated with image j (see below for details).

The benefits of this multicriterion optimization are twofold. First, automatically predicting these plaque image attributes is useful in its own right, as they appear to be clinically relevant [6]. Second, learning multiple objectives from a shared representation is known to improve the learning efficiency and prediction accuracy [20], probably by making it more difficult to overfit. However, the benefit depends on the appropriate choice of the weights w,, which are difficult and time consuming to tune manually. Fortunatelly, a maximum likelihood formulation with a Gaussian model suggested in [20] allows us to treat the weights  $w_i = 1/(2\sigma_i^2)$  as additional parameters of a modified loss

function
$$L_{mreg} = L_{reg} - 1/2 \sum_{i=1}^{A} log w_{i}$$

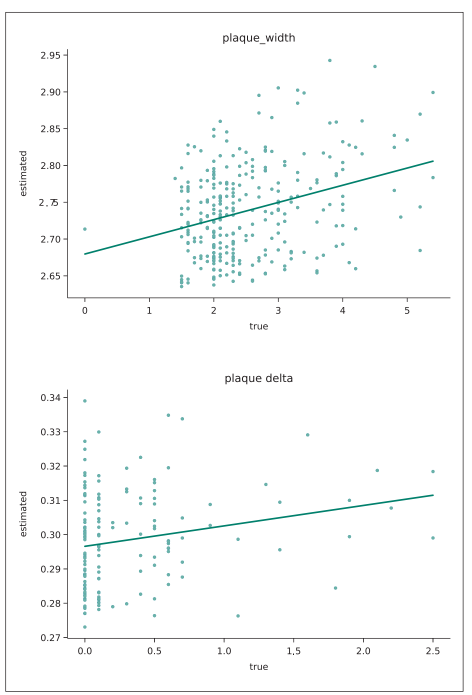


Fig. 5. Scatter plots of the estimated plaque width d (A) and the estimated increase in plaque width  $\Delta$  (B) with respect to expert annotations. The green line shows the least squares linear fit.

Obr. 5. Korelační graf odhadované šířky plátu d (A) a odhadovaného nárůstu šířky plátu  $\Delta$  (B) vzhledem k expertnímu označení. Zelená čára znázorňuje lineární aproximaci metodou nejmenších čtverců.

This way, an equilibrium is found so that well estimated attributes are given more weight. We also get an estimate  $\sigma_i^2$  of the estimation error. For numerical reasons, we optimize  $z_i = \log w_i$  instead of  $w_i$  directly.

We used a small shallow CNN [21] as the regression network R, so that it is less prone

to overfitting and concentrates on texture rather than shape. It consists of 5 convolutional layers with  $3\times3$  kernels, with 32, 64,128,256, and 64 channels, respectively, each followed by batch normalization, maxpooling, ReLU nonlinearity, and dropout. At the end, the number of channels was re-

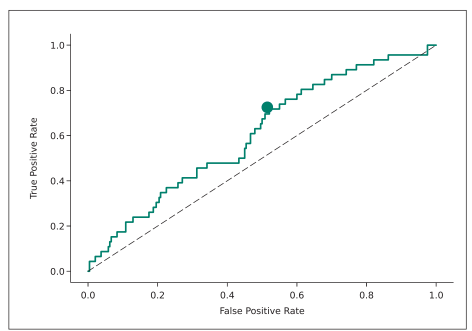


Fig. 6. ROC curve of an image-based classifier between stable and progressive plaques, defined as  $\Delta \ge 0.4$  mm. The dot corresponds to the maximum  $F_1$  working point. The dashed line corresponds to a random choice. AUC = 0.609.

AUC – area under curve; ROC – receiver operating characteristic

Obr. 6. ROC křivka klasifikátoru rozlišujícího mezi stabilními a progresivními pláty, definovaná jako  $\Delta \ge 0,4$  mm. Tečka odpovídá pracovnímu bodu maximalizujícímu hodnotu  $F_1$ . Přerušovaná čára odpovídá náhodnému výběru. AUC = 0,609.

AUC – plocha pod křivkou; ROC – receiver operating characteristic

duced to A and passed again through batch normalization, ReLU nonlinearity, dropout, and a final linear layer.

The usual approach would be to feed the regression network R the input image multiplied by a mask, leaving only the plaque region. This would allow the regression network to focus on the plaque region. Some features, like the plaque width, can be directly derived from the plaque mask. Unfortunately, this makes the regression vulnerable to segmentation errors. Alternatively, we could use the original, unmasked images as inputs, thus avoiding the dependency on the segmentation. Unfortunately, our set of image annotations was too small to train the regression network to focus on the plaque. We therefore decided to combine these approaches and use three channels as the input  $u = (u^1, u^2, u^3)$  of the regression network R. The first channel is the cropped grayscale image

$$u_i^1 = x_i$$

the second is the image limited to the plaque region using the automatic segmentation output

$$u_i^2 = x_i \left[ \underset{k}{\operatorname{argmax}} f_i^k = 3 \right]$$

where [-] is the Iverson bracket, equal to 1 if the expression is true and 0 otherwise. The third channel is the (soft) plaque mask itself

$$u_{i}^{3} = f_{i}^{3}$$

where i is the pixel index, f is the segmentation network output and 3 corresponds to the plaque class. See Fig. 4 for an example.

To encourage the network to use all channels, we randomly replaced each channel with zeros with the probability 0.5. This can be considered a channel-level dropout. Other augmentation operations were standard: random horizontal flip, rotation  $\pm$  40°, scale change  $\pm$  10%, and random cropping to the final input size 384  $\times$  384 pixels.

#### **Results**

The prediction performance was evaluated on the test part of the dataset (not used in training) by comparing the predicted and reference (expert-determined) attributes  $v_j^{\, i}$ . Table 1 shows the calculated root mean squared error (RMSE, after denormalization to the original range), the Pearson's correlation coefficient  $\rho$  and the associated probability P (significance) of an uncorrelated sys-

Tab. 2. Confusion matrix (top) and statistical performance figures (bottom) of an image-based classifier between stable and progressive plaques (defined as  $\Delta \ge 0.4 \, \text{mm}$ ) at a working point maximizing F,.

true	predicted			
liue	stable	unstable		
stable	114	121		
unstable	14	37		
AUC	0.609			
accuracy	0.528			
sensitivity	0.725			
precision	0.234			
specificity	0.485			
F <sub>1</sub>	0.354			
odds ratio	2.14			
NUC – area under curve				

tem producing |p'| > |p| under the normality assumption (calculated using the Pearson's function from the SciPy Python library). We do not show these performance criteria if the correlation is not significant at least at the 0.05 level.

We see that out of the 9 image attributes, we can predict two at the 0.01 significance level. First, the estimation of the plaque width d correlates very significantly with the reference values, although the  $\rho =$ 0.32 correlation is far from the ideal value of  $\rho = 1$ . Second, our estimate of  $\Delta$ , the future increase of the plaque width, is also significantly correlated with the reference values, although the correlation is smaller,  $\rho = 0.22$ . Fig. 5 shows the predicted and reference values as a scatter plot. We can see that the correlation is indeed present but weak. Note also that the distribution of  $\Delta$  is highly nonsymmetric – low values of  $\Delta$  are much more frequent than high values, making the regression challenging.

Besides regression, we also evaluated the prediction of  $\Delta$  in a binary classification problem framework. The two classes are stable and unstable, defined by thresholding  $\Delta \geq \tau$ . For comparability with [6,12], the threshold was set at  $\tau=0.4$  mm, which corresponds to twice the estimated plaque width measurement error. Fig. 6 shows the ROC (receiver operating characteristic curve) of this classi-

fier with a working point maximizing the  $F_1$  score. The confusion matrix and standard binary classification performance measures are shown in Tab. 2. We see that the classifier is far from perfect, but also clearly above random choice.

#### **Discussion and conclusions**

Using artificial intelligence methods for carotid plaque detection and characterization in ultrasound, as well as in other modalities, is an active area of research described in a number of review articles [22,23]. However, most of these works aimed to characterize the current state. As an example, there is a CAD (computer aided diagnostic) system [24] capable of differentiating between symptomatic and asymptomatic plaques with nearly perfect AUC = 0.956 (area under the ROC curve), with some manual steps. In that case, however, symptomatic plaque means that the patient is currently experiencing stroke, transient ischemic attack (TIA), or amaurosis fugax (AF, transient loss of vision). This is very different from the prediction task we are solving, which aims to predict future state, in order to be able to take preventive measures in time to avoid serious complications such as a stroke.

The gold standard in carotid plaque imaging is now MRI [3], with ultrasound providing much less information. A number of image features seem to be statistically relevant for distinguishing symptomatic plaques [9] but much less so for the prediction [6]. Contrast ultrasound was found not to be useful to identify vulnerable plaques in asymptomatic patients [11], which gives even less hope to non-contrast ultrasound.

Histology classification after endarterectomy was found to correlate with plaque being symptomatic [2]. There were therefore attempts to correlate the ultrasound image features with information from histology. However, the reported AUC = 0.68 (95% confidence interval 0.59–0.78) [7] is not significantly better than ours (0.609).

It can therefore be expected that predicting future plaque growth from ultrasound is difficult. Our achieved performance indicators on this task are modest but appear to be comparable to previously reported results. For example, our AUC = 0.609 (Tab. 2) is almost identical to AUC = 0.604 in [12]. Our odds ratio OR = 2.14 is very much comparable to the highest reported OR = 2.328 (for ulcerated surface, confidence interval 1.42–3.82) in [6] and higher than all

others. Let us also emphasize our relatively good performance in determining the current plaque width, which we already know to be a relevant factor for predicting the stability [6].

Since our deep-learning created features are not directly related to human-designed features used before [12], there is a hope that combining them might further improve the prediction performance. On the other hand, a pessimistic interpretation of our results is that further improvement is unlikely since the plaque growth depends on other factors that are not captured by the ultrasound image.

We yet need to explain our failure to predict the other image attributes, such as echogenicity and homogeneity. We can only hypothesize that this might be caused by inconsistent annotations, as characterizing plaques with a naked eye is often unreliable and has high intra- and inter-observer variability [22,25].

From the methodological point of view, we have applied several new techniques to both segmentation and regression, which might be useful for other tasks, too. They are based on previous work but adapted to our task. In particular, we have used GANs to model feasible segmentations, employed automatically generated weak annotations, and multi-criterion and multi-channel regression with channel-level dropout.

#### **Ethics approval**

The study was performed according to Helsinki Declaration of 1975 (and its 2004 and 2008 revisions). All participants provided an informed consent. We used data from the clinical study ANTIQUE [13], which was approved by the Ethics Board of the Faculty Hospital Ostrava (approval number 605/2014, 31 July 2014) and Ethics Committee of the Central Military Hospital Prague (approval number 108/12-44/2018, 11 June 2018). All participants provided an informed consent.

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# **Competing interests**

The authors have no relevant financial or non-financial interests to disclose.

#### **Author Contributions**

Jan Kybic performed image analysis, machine learning, implementation, experiments, writing and editing. David Pakizer, Jiří Kozel and Patricie Michalčová performed data annotation. František Charvát performed data collection and evaluation. David Školoudík suggested the concept, provided medical expertise, and worked on data acquisition, curation, and editing. All authors read and approved the final manuscript.

#### **Consent to publish**

This manuscript contains no identifiable individual's personal data.

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