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# **ORIGINAL** RESEARCH

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# 3D Quantitative Evaluation of Atherosclerotic **Plaque Based on Rotational Angiography**

BACKGROUND AND PURPOSE: Atherosclerosis is a systemic disease that has been shown to cause various cardiovascular diseases and stroke. However, technologies to evaluate the volume of atherosclerotic plaque are limited. We present a method for determination of 3D plaque volume based on RA.

MATERIALS AND METHODS: 3DRA images obtained from patients were used to evaluate the plaque. Six patients who were diagnosed with atherosclerotic lesions were included. The PR model developed for 3DRA was applied to analyze the geometry of the vessel and calculate the plaque volume. To validate the present method, we tested computer-generated phantoms with different degrees of

RESULTS: Application of PR to clinical cases allowed the estimation of plaque morphology and quantification of plaque volume. Technique validation showed that on average, PR can rebuild 92% of the plague and provide satisfactory determination of plague volume.

**CONCLUSIONS**: A new approach to obtain plaque volume based on 3DRA is presented. The initial tests in 6 clinical cases and validation with different phantoms showed that this method is feasible. Further validation in a larger clinical series is required to assess the ultimate value of the present technique.

**ABBREVIATIONS:** ACA = anterior cerebral artery; CPV = computer-estimated plague volume; ICA = internal carotid artery; MCA = middle cerebral artery; MSR = multiscale representation; PDE = partial differential equation; PR = plaque reconstruction; PV = plaque volume; RA = rotational angiography

therosclerosis is a systemic disease that has been shown to Acause various cardiovascular diseases and stroke. In particular, atherosclerotic plaque has been associated with many acute clinical complications. Cerebral infarction in the territory of the carotid arteries accounts for most strokes in Western countries and approximately 20% of patients have significant atherosclerotic narrowing at the origin of the internal carotid artery.<sup>1,2</sup> Moreover, approximately 10% of the occurrence of ischemic stroke in the United States is related to intracranial atherosclerotic disease.3,4 There is an immediate need to assess atherosclerosis in vivo to study the disease.<sup>2,5-8</sup>

Advances in imaging technology have made it possible to observe the 3D geometric changes of blood vessels and evaluate the degree of vessel narrowing in patients by using techniques such as MR imaging and CT. 6,7,9,10 Researchers have also presented various techniques and computational models to assess irregular vascular geometry and to estimate the characteristics of atherosclerotic plaque. 5,7,11-13 While plaque volume has been reported as an important tool for patient management and for evaluating new therapies, 14-16 methods to evaluate this index are limited. 14,15,17,18 Because conventional angiography remains the criterion standard for the diagnosis of vascular diseases, 19,20 techniques that can quantify plaque by using conventional angiography data can be useful.

In this article, we present a technique, PR, to quantify atherosclerotic plaque volume from 3DRA. On the basis of the

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concept of MSR,21,22 we developed the PR model to recover the vessel lumen at the regions of the atherosclerotic lesions and to compute the morphology of the plaque. Because different volumes of plaque may relate to different clinical risks, to verify that the proposed method is accurate and useful for ranges of atherosclerotic conditions, we tested and validated phantoms with various degrees of stenosis. Our initial experiences with PR in patients showed that this method can compute plaque in different arteries.

### **Materials and Methods**

# Case Selection and Imaging

Patients who were diagnosed with atherosclerosis were selected from the clinical data bases at the Division of Interventional Neuroradiology, UCLA Medical Center from 2009 to 2010. Among these patients (total of 10), 7 had 3DRA imaging of the atherosclerotic regions. One patient was excluded due to inadequate image quality for 3D reconstruction. At the end, 6 patients were selected. The locations of the atherosclerosis were the following: cases I and IV, the ACA; cases II and V, the MCA; and cases III and VI, the ICA. The 3D rotational cerebral angiograms were acquired with an Integris unit (Philips Healthcare, Best, the Netherlands). To perform the plaque computation, we transferred the 3D data to a Dell 490 workstation for the mathematic modeling and reconstruction of the plaque.

## Mathematic Modeling and Reconstruction of Atherosclerotic Plaque

Angiography captures contrast injected into the blood stream, and it is the vessel lumen that is imaged. Regions of irregular lumen in angiographic images, such as local narrowing and lack of contrast (Fig 1A), are an important indication of vascular diseases.<sup>2,5</sup> Our technique to compute the plaque is the following: First denote these geo-

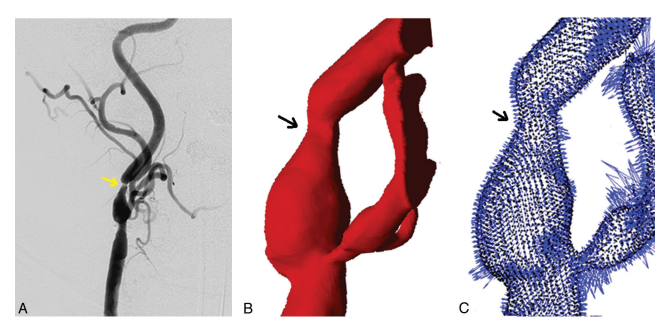


Fig 1. Representative angiographic images indicating the location of atherosclerotic plaque (yellow arrow, A) and 3D reconstruction of the same blood vessel (B) in mathematic indices ready for the plaque-estimation algorithm vector field (C), which encodes surface curvature.

metric changes with numeric indices and then extract curvature parameters to find the smooth-vessel wall.

In the PR model, the vessel geometry was first transformed into a mathematic expression (Appendix A) based on the MSR for shapes (Fig 1). $^{21,22}$  It allowed the representation of vessel geometries as a sequence of multiscaled detailed features by using level set motions. $^{23}$  Then, we implemented the fast diffusion-generated motion algorithm in the numeric procedure to solve the Hamilton-Jacobi–like equations. $^{24}$  Through the MSR, we separated the vessel detail from the smooth components of the vessel. These features were later incorporated to facilitate an accurate reconstruction of the given vessel near the atherosclerotic lesion. Finally, the vessel was recovered by filling in the data on the basis of the surrounding geometry, such as surface curvatures (Fig 1C). $^{25,26}$  At the end, by taking the difference between the reconstructed vessel and the original vessel, we could estimate the volume of plaque. Mathematic formulations and computer algorithms for the PR model are summarized in Appendices A—C.

## Technique Validation

To validate the PR algorithm, we used computer-generated phantoms created by using Matlab (MathWorks, Natick, Massachusetts) to simulate various scenarios of stenosis. Phantoms were made to represent vessels 20 mm in length and 5 mm in lumen diameter. We tested unilateral and bilateral plaque phantoms with stenoses of 30%, 60%, and 90%. For each degree of stenosis, experiments of different ranges of atherosclerotic lesions (5, 10, and 15 mm) in the direction of the vessel long axis were performed.

#### Statistical Analysis

The Wilcoxon signed rank tests were performed to analyze the results of PR. The statistical significance level was set at .05.

#### **Results**

Reconstructions of atherosclerotic plaque are shown in Fig 2 for cases I, II, and III and Fig 3 for cases IV, V, and VI. From the left to right are lesions at the ACA (Figs 2A and 3A), MCA (Figs 2B and 3B), and ICA (Figs 2C and 3C). The top row is the

diagnostic angiogram showing the atherosclerotic lesion. The second row is 3D geometry of the vessel built with RA images. The third row shows the reconstruction of the plaque for each case (yellow) and the percentage of flow obstruction by the plaque. At the bottom, the plaque subtracted from the blood vessel is presented, displaying the morphology and volume of the plaque.

#### Technique Validation

Representative phantoms with different degrees of stenosis are shown in Fig 4A. Phantoms with a smooth cylindric vessel (Fig 4B) represent healthy vessels without atherosclerotic disease. To validate the plaque quantification, we compared the CPV against the PV found by taking the difference between the healthy vessel and the vessel with stenosis. The ratio of CPV to PV for each phantom (CPV/PV) is shown in Fig 4C.

We tested cases with the plaque formed unilaterally and bilaterally with 30%, 60%, and 90% stenoses. Different lesion sizes (5, 10, and 15 mm) along the vessel axis were also included to evaluate whether PR is robust for different plaque sizes. Figure 4C shows the representative reconstruction of plaques in unilateral (left) and bilateral (right) phantoms. For the unilateral cases, on average, 92.8% of plaque (CPV/PV = 92.8%) was successfully reconstructed. There was no statistical difference between CPV and PV for 30%, 60%, and 90% unilateral stenoses (P = .109, P = .285, and P = .593, respectively). For the bilateral atherosclerotic cases, on average, 91.5% of plaque (CPV/PV = 91.5%) was successfully reconstructed. There was no statistical difference between CPV and PV for 30%, 60%, and 90% bilateral stenoses (P = .083, P =.109, and P = .109, respectively). Overall, PR was able to reconstruct 92.2% of plaque.

# **Discussion**

Conventional angiography is the standard diagnostic tool for intracranial atherosclerosis, which is often found as luminal irregularity in the angiogram, and it is assumed that the plaque

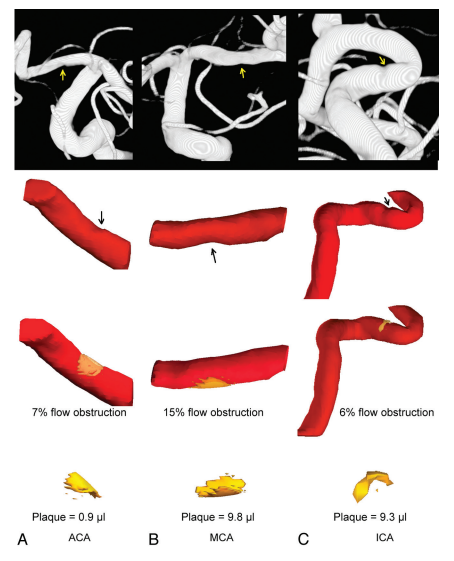


Fig 2. Plaque quantification for cases I the ACA (A), II the MCA (B), and III the ICA (C). The top row is the diagnostic images with arrows indicating stenosis. The second row shows the 3D geometry of the vessel built with 3DRA images with arrows indicating stenosis. The third row shows the reconstructed plaque (yellow) and the percentage of flow obstruction computed by the program. The bottom row shows the morphology and volume of the plaque in units of microliters.

lining the vessel wall results in such irregularity. <sup>19,20,27</sup> The purpose of this study was to develop a technique that can evaluate the volume of plaque by using conventional angiography. We introduced the PR model to reconstruct plaque on the basis of MSR principles that have been applied to study various biologic shapes. <sup>21,22</sup>

There has been much effort to develop evaluation criteria and measurement methods to study atherosclerotic plaque. Researchers have shown that the plaque volume can be a reliable measure for studying atherosclerosis. A recent randomized trial also used this quantity as 1 of the indices to track the plaque changes with time. Therefore, there is considerable value in quantifying the plaque volume from 3D angiograms. Our preliminary results showed that using PR to assess plaque volume is feasible for clinical images. Because an atherosclerotic lesion is often diagnosed or verified with catheter angiograms, this technique can help to estimate the morphology and volume of the plaque and to study the disease.

In the technique validation, we performed tests by using

phantoms with different degrees of stenosis. The results showed that the algorithms allow the reconstruction of plaque in different scenarios. On average, 92% of the plaque can be rebuilt through the proposed method, regardless of the lesion size. Additional effort such as implementing advanced computation schemes can be helpful to improve the CPV/PV ratio of the current model. Furthermore, research using postsurgical specimens to validate the plaque volume estimation is essential to understand the accuracy of the present method.

#### **Conclusions**

We present a new approach to estimate the volume of atherosclerotic plaque on the basis of 3DRA. Analysis and validation with different phantoms to simulate various degrees of stenosis have shown that this method provides satisfactory volume computation. The initial results of the clinical application suggest that this method is feasible. Further validation in a larger clinical series is required to assess the ultimate value of the present technique.

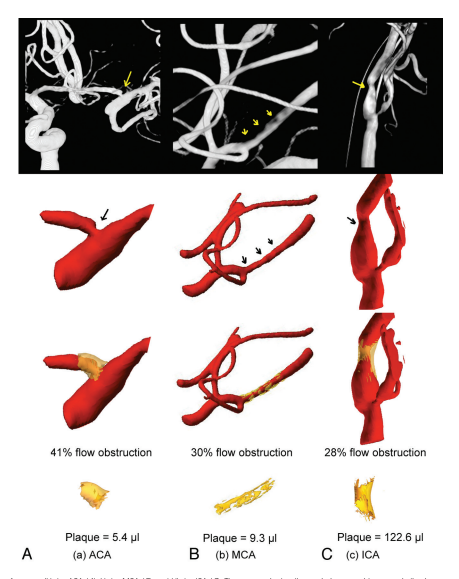


Fig 3. Plaque quantification for cases IV the ACA (A), V the MCA (B), and VI the ICA (C). The top row is the diagnostic images with arrows indicating stenosis. The second row shows the 3D geometry of the vessel built with 3DRA images and arrows indicating stenosis. The third row shows the reconstructed plaque (yellow) and the percentage of flow obstruction. The bottom row shows the morphology and volume of the plaque in units of microliters.

#### **Appendix**

#### A) Vessel Lumen Geometry

We first reconstructed the blood vessel lumen and expressed boundaries  $\partial D$  of domains  $D \in \mathbb{R}^3$  by level set functions<sup>23</sup>  $\phi$  as the following:

1) 
$$\phi(x) \begin{cases} <0, & x \in D \\ >0, & x \in D^c \end{cases}$$

The multiscale representation of a blood vessel then can be obtained by level set motions through the following Hamilton-Jacobi–like equation:

2) 
$$\phi_t + v_n(\nabla \phi) |\nabla \phi| = 0 \quad \phi(x, 0) = \phi_0(x),$$

where  $\phi_0$  is the level set function of the given blood vessel,  $v_n$  is the velocity field, and  $t \in [0, T]$ . To obtain a second-order approximation to PDE (equation 2) and reduce the complexity of solving the nonlinear PDE, the diffusion-generated motion technique was used to solve PDE (equation 2).<sup>24</sup> Therefore, the continuous transform of the vessel  $\phi_0$ , and its inverse

transform can be expressed as equations 3 and 4 respectively with  $\phi(x, t)$ , the solution of PDE (equation 2):

3) 
$$W(\phi_0) := \overrightarrow{W}(x, t) := - \nu_n \frac{\nabla \phi}{|\nabla \phi|}$$

4) 
$$\psi_t + \overrightarrow{W}(x, t) \cdot \nabla \psi = 0, \quad \psi(x, 0) = \phi(x, T).$$

Then, we denote  $\overrightarrow{W}_1(x, t)$  as the restriction of  $\overrightarrow{W}(x, t)$  on  $S_t := \{x: \phi(x, t) = 0\}$ , and express an MSR for the original vessel  $S_o$  as equation 5, which is invariant under rigid-body transformation and initial embeddings of  $\phi_0$ :

5) 
$$MSR(S_0) = \{\{\overrightarrow{W}_1(x,t)\}_{t \in [0,T]}, S_0\},$$

#### B) Reconstruction of Atherosclerotic Plaque

We used MSR as described above for the geometric representations of the blood vessel, where we chose the normal-velocity field in PDE (equation 2) to be  $v_n = c + \kappa_a - \kappa$  (Fig 1*C*).  $\kappa$  and  $\kappa_a$  are the mean curvature and average mean curvature, respectively. c is constant. After solving PDE (equation 2)<sup>24</sup>,

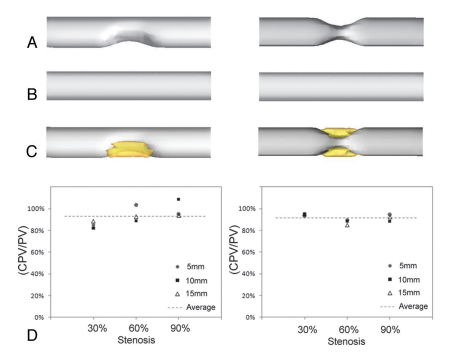


Fig 4. A, Representative unilateral (left) and bilateral (right) stenosis phantoms. B, The ideal vessel without stenosis. C, The reconstructed plaque for the unilateral (left) and bilateral (right) models. D, The percentage of the volume estimation (CPV/PV) for lesion sizes 5, 10, and 15 mm in the direction of the vessel long axis.

we recorded the details  $\overrightarrow{W}(x,t)$  for all x in a narrow band of  $\phi(x,t)$ . This step is crucial to ensure that the reconstruction is accurate. Finally, the vessel was reconstructed with  $\psi_t + \overrightarrow{W}(x,t) \cdot \nabla \psi = \epsilon \nabla_2 \psi$ , where  $\epsilon$  is diminished for each iteration. When  $\epsilon$  goes to zero, the solution will converge to the solution of equation 4. Therefore, a diminishing  $\epsilon$  ensures the final convergence of our calculation.

# C) Summary of Algorithms to Reconstruct Atherosclerotic Plaques

1) Perform MSR transform of the given vessel by solving PDE (2) with

$$v_n = c + \kappa_a - \kappa$$

by using the fast diffusion-based level set motion.

- 2) Record the details  $\widetilde{W}(x,t)$  for all x in a narrow band of  $\phi(x,t)$ .
- 3) Reconstruct the blood vessel through the following equations:

$$\psi_i + \overrightarrow{W}(x, t) \cdot \nabla \psi = \epsilon \nabla^2 \psi, \quad \psi(x, 0) = \phi(x, T).$$

4) Apply the data outside the inpainting domain and repeat until the level set function satisfies the following equation:

$$\|\phi \ current(x,0) - \phi \ previous(x,0)\| < 0.01$$

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