

# Vesselness-guided Active Contour: A Coronary Vessel Extraction Method

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#### **ABSTRACT**

Vessel extraction is a critical task in clinical practice. In this paper, we propose a new approach for vessel extraction using an active contour model by defining a novel vesselness-based term, based on accurate analysis of the vessel structure in the image. To achieve the novel term, a simple and fast directional filter bank is proposed, which does not employ down sampling and resampling used in earlier versions of directional filter banks. The proposed model not only preserves the performance of the existing models on images with intensity inhomogeneity, but also overcomes their inability both to segment low contrast vessels and to omit non-vessel structures. Experimental results for synthetic images and coronary X-ray angiograms show desirable performance of our model.

Key words: Active contour, vesselness-based, vessel extraction

# INTRODUCTION

Vessel extraction is a critical task in clinical practice. Correct assessment, especially accurate visualization and quantification of blood vessels traced in angiograms plays a significant role in a number of clinical diagnostic procedures, e.g. diagnosis relied on vessel width, reflectivity, tortuosity and abnormal branching. For example, appearance of vessels of a certain width in the heart may reveal the signs of stenoses. Grading of stenoses is of importance to diagnose the extent of vascular disease which determines the treatment therapy.<sup>[1,2]</sup> Although it is possible for medical experts to delineate vessels, but manual delineation of the vasculature becomes tedious or even impossible when the number of vessels in an image is large, or when a large number of images have to be inspected. Therefore, the development of automatic and accurate vessel-tree reconstruction from angiograms is highly desirable. In recent years, this has been a challenging task. The key fact is that vessels cannot be characterized uniformly. Since the blood, either by itself or by the contrast agent injected into it, is responsible for the vessel contrast to the background, vessels with larger widths usually have high contrast while smaller ones resemble the background. Many segmentation methods have been used to visualize the blood vessel structures in the human body. A review of vessel extraction techniques and algorithms can be found in. [3] A combination of Hessian matrix multi-scale filtering and region-growing is used to segment the coronary artery in angiograms.[4] An automatic technique, which provides correspondence between extracted vessels and pre-procedural vessels, improves the quality of disease diagnosis using multiple images and follow-up and provides important information for non-rigid image registration.[5] A segmentation method based on fusion algorithm is proposed in. [6] In this method, firstly, the coronary arteries are extracted by using the maximizing entropy segmentation method based on top-hat. Then the coronary arteries are extracted again by using the maximizing entropy segmentation method based on Gaussian filter. Finally, the last segmentation result is the image obtained by fusing two extracted coronary arteries images. A region-growing method is proposed for vessel segmentation in angiography images. The method consists of two parts: The feature map extraction based on a novel vesselness function; and the segmentation process which includes automatic seed-point selection, main branch segmentation and vessel detail repair. Both the greyscale and spatial information are extracted for segmentation based on region growing algorithm.<sup>[7]</sup> Furthermore, a classification approach is proposed to distinguish between the coronary arteries and background images using the shape context descriptor and the learning framework of spatial pyramid kernels. [8] In this paper, we focus on the new development of angiograms segmentation based on an active contour model.

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Active contour models are among the most successful image segmentation techniques in clinical applications. Active contours may be categorized as edge based<sup>[9-11]</sup> or region based.<sup>[12-15]</sup>

In region-based models, the statistical information inside and outside the contour are used to control the evolution. They are less sensitive to noise and are more effective for images with weak edges or images without edges. However, popular region-based active contour models<sup>[12,13]</sup> and<sup>[14]</sup> tend to rely on intensity homogeneity of each region to be segmented. For example, the popular piecewise constant models are based on the assumption that image intensities are statistically homogeneous (roughly constant) in each region.

In fact, intensity inhomogeneity often occurs in real images from different modalities. Generally for medical images, inhomogeneity is usually due to technical limitations or artifacts introduced by the object being imaged. In a study done by Li *et al.*,<sup>116</sup> Zhang *et al.*<sup>117</sup> and Wang *et al.*<sup>118</sup> proposed local binary fitting (LBF), local image fitting (LIF) and local Gaussian distribution (LGD) fitting models, respectively, capable to segment images with intensity inhomogeneity by embedding the local image information. When intensity inhomogeneity occurs in angiograms, each of these models can be used to segment the vessels. LIF model is more computationally efficient than LBF, but it cannot accurately extract vessels. Therefore, LBF model is more suitable for vessel segmentation in angiograms but it may fail to extract weak vessels.

In this paper, we propose a new approach for vessel extraction using LBF model by defining a novel vesselness-based term which segments low contrast vessels and omits non-vessel structures. The results are compared with the 4 active contour models.<sup>[12,16-18]</sup>

The remainder of this paper is organized as follows: In section 2, the LBF model is described. The proposed method is stated in section 3. Experimental results on synthetic images and real angiograms are presented in section 4 and section 5 is devoted to conclusion.

# LBF MODEL

Let  $\Omega \subset \mathbb{R}^2$  be the image domain and  $I: \Omega \to \mathbb{R}$  be a given gray level image. To segment the images with intensity inhomogeneity, in  $| ^{16} |$  a region-based model using intensity information inlocal regions at a controllable scale is proposed. For an image, I(x, y) on the image domain  $\Omega$ , the algorithm proposes to minimize the following region-scalable fitting energy of a contour C:

$$\begin{split} F_{1}(C, f_{1}(P), f_{2}(P)) &= \int_{\Omega} [\lambda_{1} \int_{\Omega_{1}} K_{\sigma}(P-Q) | I(Q) - f_{1}(P) |^{2} dQ \\ &+ \lambda_{2} \int_{\Omega_{2}} K_{\sigma}(P-Q) | I(Q) - f_{2}(P) |^{2} dQ] dP \end{split} \tag{1}$$

where, P=(x,y),  $\lambda_1$  and  $\lambda_2$  are positive constants, C is the contour,  $f_1(P)$  and  $f_2(P)$  are two values that approximate image intensities in  $\Omega_1$  and  $\Omega_2$  (regions inside and outside the contour C, respectively),  $K_{\sigma}$  is a Gaussian kernel with a standard deviation  $\sigma$ . I(Q) is the intensity of pixels in a window around P.

The energy in (1) is incorporated into a level set formulation. In the level set method, the curve  $C \subset \Omega$  is represented by the zero level set of Lipschitz function $\phi$ :  $\Omega \subset R$ . Minimizing the energy functional  $F_I$  with respect to  $\phi$  using the gradient descent method leads to:

$$\frac{\partial \phi}{\partial t} = -\delta_{\varepsilon}(\phi)(\lambda_1 e_1 - \lambda_2 e_2) \tag{2}$$

Where  $e_1$  and  $e_2$  are the functions

$$e_i(P) = \int K_{\sigma} (Q - P) |I(P) - f_i(Q)|^2 dQ, i = 1, 2$$
 (3)

with

$$f_{1}(P) = \frac{K_{\sigma}(P) * [H_{\varepsilon}(\phi(P))I(P)]}{K_{\sigma}(P) * H\varepsilon(\phi(P))}$$

$$f_{2}(P) = \frac{K_{\sigma}(P) * [(1 - H_{\varepsilon}(\phi(P)))I(P)]}{K_{\sigma}(P) * (1 - H\varepsilon(\phi(P)))}$$
(4)

In practice, the Heaviside function in the above energy functional is approximated by a smooth function defined by

$$H_{\varepsilon}(x) = \frac{1}{2} \left[ 1 + \frac{2}{\pi} \arctan(\frac{x}{\varepsilon}) \right]$$
 (5)

Hence

$$\delta_{\varepsilon}(x) = H_{\varepsilon}'(x) = \frac{1}{\pi} \frac{\varepsilon}{\varepsilon^2 + x^2}$$
 (6)

As proposed in,<sup>[11]</sup> in order to maintain the regularity of the level set function a regularization term is incorporated into (2). Moreover, to regularize the zero level contour, the arc length term is also added to the energy function (2). After all, the level set evolution equation becomes:

$$\begin{split} \frac{\partial \phi}{\partial t} &= -\delta_{\varepsilon}(\phi)((\lambda_{1}e_{1} - \lambda_{2}e_{2}) \\ &+ \upsilon(\nabla^{2}\phi - div(\frac{\nabla\phi}{|\nabla\phi|})) + \mu \ \delta_{\varepsilon}(\phi)div(\frac{\nabla\phi}{|\nabla\phi|}) \end{split} \tag{7}$$

LBF model of (7) can well handle image with intensity inhomogeneity and is proper for angiograms segmentation.

Since the standard deviation,  $\sigma$  of the kernel controls the region-scalability,<sup>[16]</sup> for better results in angiograms,  $\sigma$  should be chosen according to size of vessels. In these images, a vessel tree is composed of two main parts:

The wide vessels and the thin ones. However in LBF model, only one scale parameter was used and hence that  $\sigma$  should be proportional to wide or thin vessels. If  $\sigma$  is appropriate for small vessels, it may cause undesirable results regarding low contrast vessels. Therefore,  $\sigma$  should be chosen proportional to wide vessels, making thin ones non-extractable. On the other hand, the statistical properties of these vessels also are different and the application of the model in (7) alone fails to extract all vessels. To address these problems, we add an energy term to LBF model as described in the following.

# PROPOSED METHOD

#### Vesselness

In general, in the literature vessel enhancement in angiograms are Hessian-based filters, which are found to be sensitive to noise and sometimes give discontinued vessels due to junction suppression. To overcome these drawbacks, Truc  $et\ al.^{[19]}$  proposed the use of line-like directional features of the image to obtain more precise Hessian analysis in noisy environments and to avoid junction suppression. The features are extracted by decimation-free directional filter bank (DDFB) which decomposed the input image to a set of directional images. Thus, it can correctly reveal small and thin vessels and yield the continuous vessel tree. Accordingly, in this paper we use this method to obtain enhanced image  $D\ (Q)$  needed in (10) except that we proposed a simple and fast method of the filter bank construction named fast directional filter bank (FDFB) presented in the following.

#### **FDFB**

In order to construct the directional filter bank (DFB) or DDFB,<sup>[19]</sup> it takes three stages. Through these stages, we have to modulate, down sample, resample and post-sample a pair of diamond-shaped like passband filters in DFB (or hourglass-shaped like passbands filters in DDFB). However in the FDFB directly and easily we partition the frequency domain using one stage division algorithm.

To achieve FDFB, we consider the frequency domain in polar co-ordinates as shown in Figure 1. It is supposed that the spectrum is divided into n parts, the i<sup>th</sup> part is between  $\theta_{i,min}$  and  $\theta_{i,max}$  and the center of the i<sup>th</sup> part is

$$\theta_i = \frac{\theta_{i,\min} + \theta_{i,\max}}{2} \tag{8}$$

A mathematical description of FDFB for  $2^k$  DFBs is given by (9):

$$F_{n} = \begin{cases} 1, & \text{for } \begin{cases} \frac{(n-1)\pi}{2^{k}} \le \theta < \frac{n\pi}{2^{k}} \\ \forall r \text{ with } 0 \le u, v \le M \end{cases} \\ 0, & \text{Otherwise} \end{cases}$$
 (9)

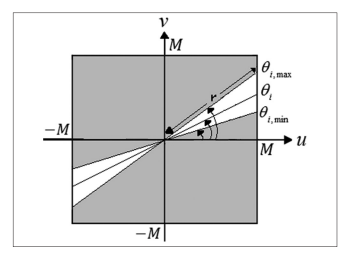


Figure 1: Consideration of frequency domain in polar coordinates

In which, u and v are frequency,  $n \in \{1, 2,..., 2^k\}$ ,  $r = \sqrt{u^2 + v^2}$  and  $\theta = \arctan(v/v)$ . The ultimate partitioning of frequency domain in FDFB is equivalent to the output partitioning of DDFB, but it competes with less computational complexity.

One major problem with DDFB and FDFB is the inevitable ringing artifact. One way to remove ringing artifact is to smooth the filters transition band with a proper low pass Gaussian filter. Smoothed filter bank for k=3 is shown in Figure 2. This filter bank partitions the frequency domain into eight sub-bands. In order to prevent aliasing, the input image is zero-padded to two times of its size. Since the angiographic images are square, we suppose the input image is  $M \times M$ . Hence, the frequency representation will be of the size  $2M \times 2M$ . The result of vessel enhancement in angiograms by Hessian-based filters is shown in Figure 3. As it demonstrates, some non-vessel parts of the image are enhanced.

# **Two-stage Level Set Formulation**

Our method combines the two energy function based on (1) as:

$$F(C) = \alpha_1 F_1(C, f_1, f_2) + \alpha_D F_D(C, d_1, d_2)$$
(10)

Where energies  $F_{i}$  and  $F_{D}$  are defined:

$$F_{I}(\phi, f_{1}, f_{2}) = \int \left(\sum_{i=1}^{2} \lambda_{i} \int_{\Omega_{i}} K_{\sigma_{0}}(P - Q)I(Q) - f_{1}(P)\right)^{2} M_{i}^{\varepsilon}(\phi(Q))dQ)dP$$
 (11)

And

$$F_{D}(\phi, d_{1}, d_{2})$$

$$= \int \left(\sum_{i=1}^{2} \lambda_{d_{i}} \int_{\Omega_{i}} K_{\sigma_{1}}(P-Q) \left| D(Q) - d_{i}(P) \right|^{2} M_{i}^{\varepsilon}(\phi(Q)) dQ \right) dP \quad (12)$$

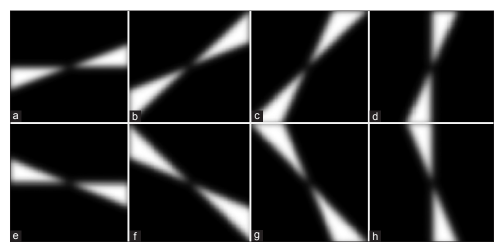


Figure 2: Partitioning of frequency domain, after smoothing filter is applied

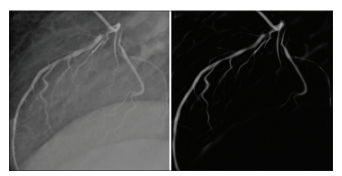


Figure 3: The result of vessel enhancement in angiograms by Hessian-based filters

In the first term of (10), I (Q) is the intensity of pixels in the original image and  $\sigma_0$  is proportional to width of wide vessels. In the second term D (Q) is the intensity of pixels in enhanced image obtained by FDFB-based method (as presented in section 3.1) and includes the vessel structure.  $d_{\tau}$  (P) and  $d_{\tau}$  (P) are two values that approximate enhanced image intensities in  $\Omega_{\tau}$  and  $\Omega_{\tau}$  (regions inside and outside the contour C, respectively) and  $\sigma_{\tau}$  is proportional to the width of thin vessels.

Finally, it is proposed to minimize the energy functional,

$$W(\phi) = \alpha_1(F_1(\phi, f_1, f_2) + \nu \vartheta(\phi)) + \alpha_p F_p(\phi, d_1, d_2) + \mu \rho(\phi)$$
 (13)

Where  $\alpha_{p}$ ,  $\alpha_{p}$ , v,  $\mu$  are regulating parameters and  $\rho$  (|*varphi*) is a level set regularization term as presented in<sup>[11]</sup>

$$\rho(\phi) = \int (|\nabla \phi(P)| - 1)^2 dP \tag{14}$$

Also, the arc length term is:

$$\vartheta(\phi) = \int_{\Omega} |\nabla H_{\varepsilon}(\phi(P))| dP \tag{15}$$

Let H be smoothed Heaviside function as defined in (5), then the proposed energy function  $W(\phi)$  can be rewritten as (16)

$$W(\phi) = \alpha_{I} \left[ \int \left( \sum_{i=1}^{2} \lambda_{i} \int_{\Omega_{i}} K_{\sigma_{0}}(P - Q) \left| I(Q) - f_{i}(P) \right|^{2} M_{i}^{\varepsilon}(\phi(Q)) dQ \right) dP \right]$$

$$+ \upsilon \int_{\Omega} \delta_{\varepsilon}(P) \left| \nabla \phi(P) \right| dP \right] + \mu \int_{\Omega} \frac{1}{2} (\left| \nabla \phi(P) \right| - 1)^{2} dP$$

$$+ \alpha_{D} \left[ \int \left( \sum_{i=1}^{2} \lambda_{di} \int_{\Omega_{i}} K_{\sigma_{1}}(P - Q) \left| D(Q) - d_{i}(P) \right|^{2} M_{i}^{\varepsilon}(\phi(Q)) dQ \right) dP \right]$$
(16)

where  $M_1^{\varepsilon}(\phi) = H_{\varepsilon}(\phi)$ ,  $M_2^{\varepsilon}(\phi) = 1 - H_{\varepsilon}(\phi)$ .

According to  $f_1$  the functions  $f_1$  (P) and  $f_2$  (P) satisfy the following Euler–Lagrange equations:

$$\int K_{\sigma_0}(P-Q)M_i^{\varepsilon}(\phi(Q)(I(Q)-f_i(P))dQ = 0, i = 1.2$$
(17)

From (17)

$$f_i(P) = \frac{K_{\sigma_0}(P) * \left[ M_i^{\varepsilon}(\phi(P))I(P) \right]}{K_{\sigma_0}(P) * M_i^{\varepsilon}(\phi(P))}, i = 1, 2$$
(18)

And  $d_1\left(P\right)$  and  $d_2\left(P\right)$  satisfy the following Euler–Lagrange equations:

$$\int K_{\sigma_{1}}(P-Q)M_{i}^{\varepsilon}(\phi(Q)(D(Q)-d_{i}(P))dQ = 0, \ i = 1, 2$$
(19)

From (19):

$$d_{i}(P) = \frac{K_{\sigma_{1}}(P) * \left\lfloor M_{i}^{\mathcal{E}}(\phi(P))D(P) \right\rfloor}{K_{\sigma_{1}}(P) * M_{i}^{\mathcal{E}}(\phi(P))}, i = 1, 2$$
(20)

By keeping  $f_1(P)$ ,  $f_2(P)$  and  $d_1(P)$ ,  $d_1(P)$  fixed and minimizing the energy function  $W(\phi)$ , <sup>th</sup>e level set evolution equation becomes:

$$\frac{\partial \phi}{\partial t} = \delta_{\varepsilon}(\phi) \left\{ \alpha_{I} \left[ \upsilon \operatorname{div}(\frac{\nabla \phi}{|\nabla \phi|}) - (\lambda_{1}e_{1} - \lambda_{2}e_{2}) \right] + \alpha_{D} \left[ (\lambda_{d1}e_{d1} - \lambda_{d2}e_{d2}) \right] \right\} + \mu \left( \nabla^{2}\phi - \operatorname{div}(\frac{\nabla \phi}{|\nabla \phi|}) \right) \tag{21}$$

Where  $e_1$  and  $e_2$  are the functions

$$e_i(P) = \int K_{\sigma_0}(Q - P) |I(P) - f_i(Q)|^2 dQ, i = 1, 2$$
 (22)

 $e_{d_1}$  and  $e_{d_2}$  are the functions.

$$e_{d_i}(P) = \int K_{\sigma_i}(Q - P) |D(P) - d_i(Q)|^2 dQ, i = 1, 2$$
 (23)

# IMPLEMENTATION AND EXPERIMENTAL RESULTS

# **Implementation**

In a Core2 Duo CPU, 2.2 GHz and 4 GB RAM, it takes about 18 s for DDFB to create filter bank and for FDFB it takes about 3 s.

In our approach, the level set function (21) was evaluated in two stages. In the first stage, in order to propagate the contour to the main vessels,  $\alpha_{\rm l}$ ,  $\alpha_{\rm p}$  are set to one and zero, respectively and  $\lambda_{\rm l}=\lambda_{\rm 2}=1$ ,  $\mu=1$ ,  $\upsilon=65$ ,  $\Delta t=0.1$ , the same as. In the second stage to segment thin vessels,  $\alpha_{\rm l}$ ,  $\alpha_{\rm p}$  are set to zero and 1, respectively and  $\lambda_{\rm dl}=\lambda_{\rm d2}=1$ . It should be mentioned that the second stage is only effective in the narrowband case.

In the first term of (21),  $\sigma_0$  is selected according to the width of main vessels and in the second term,  $\sigma_1$  is selected proportional to width of thinner vessels. For angiograms  $\sigma_0 = 3$ ,  $\sigma_1 = 1$  is desirable.

# **RESULTS**

To evaluate the result of the proposed algorithm for vessel segmentation in angiograms, it has been tested with synthetic and real images.

Figure 4 shows two synthetic images corrupted by intensity inhomogeneity. The image in Figure 4a is a synthetic image convolved with a Gaussian function to model the real angiogram. The results of the original image segmentation by LBF model and our method are shown in Figure 4b and c, respectively.

The image in Figure 4d is generated by adding Gaussian noise with  $\sigma = 0.005$  to Figure 4a. The segmentation result for noisy image by LBF model and our method are shown at Figure 4e and f, respectively.

It is apparent that our algorithm segments the vessels more precisely than LBF in noisy images. Also, the results of our algorithm for the original image and the image with a high level of noise are very close as shown in Figure 4e.

Figure 5 compares the performance of our proposed algorithm on two real coronary X-ray angiograms. In the left most column, the original images are of size 300 × 300. The second and third columns show the results of converged contour of LBF and our method for real coronary X-ray angiograms. The fourth and fifth columns show the results of segmentation of LBF and proposed method respectively. It is clear that our algorithm has been able to extract vessels with weak boundaries more precisely. In addition, our algorithm demonstrates the width of the main vessels more accurate than that of LBF.

As shown in Figure 6, the results of LBF and our method on another angiogram are shown. Our method extracts the vessels in angiograms with a complex background and noise, accurately.

In order to compare our results with the results of the four above mentioned active contour models, we randomly selected 15 X-ray coronary angiograms obtained in Sina Heart Hospital, Isfahan, Iran. These images were segmented manually by an expert at this center as a gold standard.

For performance comparison, we have selected the segmentation accuracy (ACC) measure, determined by true positive fraction (TPF) and the false positive fraction (FPF). The parameter TPF, also called "sensitivity," is the ratio of the number of pixels correctly classified as vessel pixels true positive (TP), to the total number of vessel pixels in the gold standard segmentation,

$$TPF = \frac{TP}{P} = \frac{TP}{TP + FN} = sensitivity$$

where FN is the number of pixels incorrectly classified as non-vessel pixels. The ideal amount of TPF is equal to 1. The parameter FPF is the number of pixels incorrectly classified as vessel pixels false positive, divided by the total number of non-vessel pixels in the gold standard:

$$FPF = \frac{FP}{N} = \frac{FP}{FP + TN} = 1$$
-specificity

Here, TN is the number of pixels correctly classified as non-vessel pixels. The ideal amount of TPF is equal to 0. The ACC for one image is the ratio of the total number of correctly classified points (sum of TPs and true negatives) by the total number of points in the image.

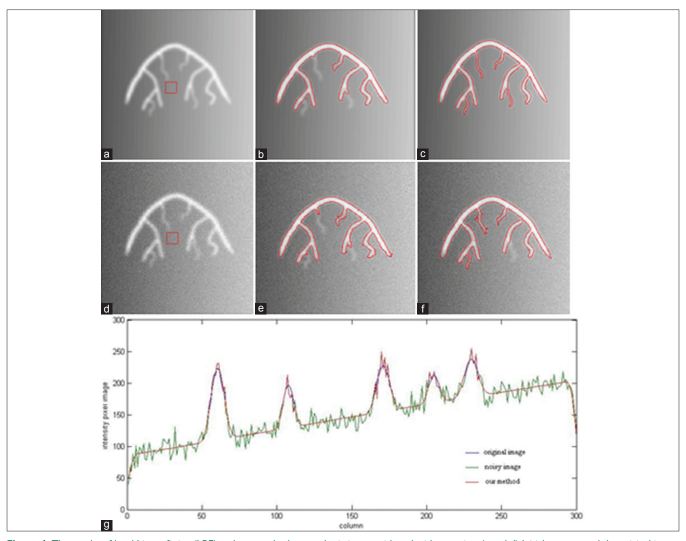


Figure 4: The results of local binary fitting (LBF) and our method on synthetic image with and without noise: (a and d) Initial contour and the original image (b and c) Final contour of clean image by LBF model and our method, respectively (e and f) Final contour of noisy image by LBF model and our method, respectively (g) Cross-section of the original image (blue line), noisy image (green line) and our method (red line)

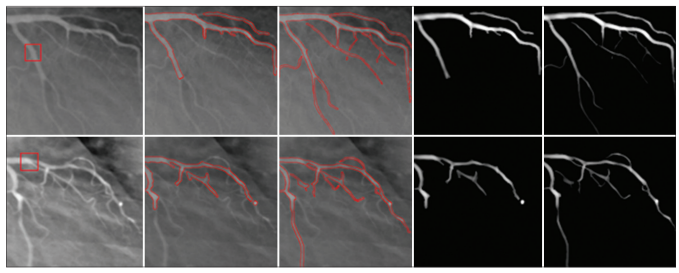


Figure 5: The results of local binary fitting (LBF) model and our method for coronary X-ray angiograms. Original images and initial contours (first column), final contour by LBF method (second column), our method (third column), segmentation result by LBF method (forth column) and by our method (fifth column)

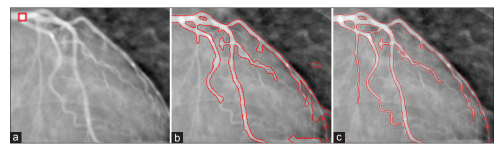


Figure 6: The results of local binary fitting (LBF) and our method on an angiogram. (a) Original image and initial contour. (b) LBF. (c) Our method

Table 1: Performance of vessel segmentation by active contour methods

	True positive fraction	False positive fraction	Accuracy (standard deviation)
Our method	0.8943	0.012	0.9747 (0.0085)
LBF <sup>[11]</sup>	0.8127	0.018	0.9053 (0.0102)
LIF <sup>[12]</sup>	0.1812	0.008	0.7524 (0.0072)
LGD <sup>[13]</sup>	0.3012	0.025	0.8471 (0.0192)
Chan and Vese <sup>[7]</sup>	0.1732	0.2326	0.6596 (0.0135)

LBF – Local binary fitting; LIF – Local image fitting; LGD – Local Gaussian distribution

$$ACC = \frac{TP + TN}{P + N} = \frac{TP + TN}{TP + FN + FP + TN}$$

The ideal amount of ACC is also equal to 1.

Table 1 shows the average of the ACC measurements for all 15 angiograms for our method, LBF, LIF, LGD and Chan-Vese models. It is apparent that the performance of our method is better than the other methods.

# **CONCLUSIONS**

In this paper, we have presented a new active contour model for vessel segmentation presented in (21). In the level set formulation, the new approach combines both intensity information in regions with a controllable scale and information of the image enhanced by FDFB-based method. The first term in (21) extracts the main vessels using LBF model, by incorporating the vessel structure while the second term extracts thin vessels. Applying LBF model alone fails to extract all vessels. Also, if the first term is omitted, the vesselness-based term might segment non-vessel structures. Therefore, a combination of the two terms leads to a better result than using each term alone. The model is efficient for the segmentation of vessels as well as other line-like structures. In conclusion, the proposed method is a more accurate candidate for segmentation coronary vessels used in clinical tasks.

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