

● *Original Contribution*

## INTRALUMINAL ULTRASONIC PALPATION IMAGING TECHNIQUE REVISITED FOR ANISOTROPIC CHARACTERIZATION OF HEALTHY AND ATHEROSCLEROTIC CORONARY ARTERIES: A FEASIBILITY STUDY

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**Abstract**—Accurate mechanical characterization of coronary atherosclerotic lesions remains essential for the *in vivo* detection of vulnerable plaques. Using intravascular ultrasound strain measurements and based on the mechanical response of a circular and concentric vascular model, E. I. Céspedes, C. L. de Korte and A. F. van der Steen developed an elasticity-palpography technique in 2000 to estimate the apparent stress–strain modulus palpogram of the thick subendoluminal arterial wall layer. More recently, this approach was improved by our group to consider the real anatomic shape of the vulnerable plaque. Even though these two studies highlighted original and promising approaches for improving the detection of vulnerable plaques, they did not overcome a main limitation related to the anisotropic mechanical behavior of the vascular tissue. The present study was therefore designed to extend these previous approaches by considering the orthotropic mechanical properties of the arterial wall and lesion constituents. Based on the continuum mechanics theory prescribing the strain field, an elastic anisotropy index was defined. This new anisotropic elasticity-palpography technique was successfully applied to characterize ten coronary plaque and one healthy vessel geometries of patients imaged *in vivo* with intravascular ultrasound. The results revealed that the anisotropy index-palpograms were estimated with a good accuracy (with a mean relative error of  $26.8 \pm 48.8\%$ ) compared with ground true solutions. (E-mail: [jacques.ohayon@univ-grenoble-alpes.fr](mailto:jacques.ohayon@univ-grenoble-alpes.fr)) Crown Copyright © 2018 Published by Elsevier Inc. on behalf of World Federation for Ultrasound in Medicine & Biology. All rights reserved.

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### INTRODUCTION

Morphologic and mechanical characterization of coronary atherosclerotic lesions remains an essential step for the diagnosis of rupture-prone plaques and subsequent thrombosis (Alsheikh-Ali et al. 2010; Arbab-Zadeh et al. 2012; Benjamin et al. 2017; Fleg et al. 2012; Libby et al. 2002; Naghavi et al. 2003).

To detect vulnerable plaques (VPs) in a timely manner and prevent adverse coronary events, several endovascular imaging techniques are available to clinicians or in development including intravascular ultrasound (IVUS) (Cheng et al. 2014; Rioufol et al. 2002), optical coherence tomography (Sinclair et al. 2015), magnetic resonance imaging (Cavalcante and Larose 2016; Larose et al. 2005) and near-infrared spectroscopy (Caplan et al. 2006; Madder et al. 2016), as well as emergent hybrid techniques that combine two or more imaging modalities (Abran et al. 2014; Bourantas et al.

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2017). Despite these technological advances, identifying which VPs will rupture and induce an acute event remains a challenging task.

Previous biomechanical studies have identified peak cap stress amplitude as a strong indicator of plaque vulnerability to rupture (Finet et al. 2004; Gijssen and Migliavacca 2014; Kok et al. 2016; Ohayon et al. 2008; Tang et al. 2009). Others have established that plaque rupture risk is also attributed to additional factors such as the presence of microcalcifications in the thin cap (Cardoso and Weinbaum 2014; Kelly-Arnold et al. 2013; Thondapu et al. 2017), plaque morphology (Akyildiz et al. 2016; Ohayon et al. 2008) and hemodynamics (Ahmadi et al. 2016; Gijssen et al. 2008; Morbiducci et al. 2016), as well as tissue mechanical properties of the atherosclerotic lesion (Guo et al. 2017; Finet et al. 2004; Ohayon et al. 2014).

Although plaque morphology and cap thickness can be directly extracted from endovascular images, the stress distribution within the plaque is impossible to obtain using current medical imaging technologies. However, the local deformation (strain) of the arterial wall and plaque caused by the cyclic intraluminal pressure can be calculated from a series of ultrasound images (Doyley 2012; Golemati et al. 2016; Maurice et al. 2004b). Based on the postulate that local variations in elastic properties of tissues are generally correlated with pathologic regions, strain elastography—a strain field-based technique—was developed. This approach was first introduced by Ophir's group (Céspedes et al. 1993; Ophir et al. 1991) to map the mechanical strain response to an applied force. Nevertheless, complex non-linear geometric effects do not allow a direct correlation between the measured spatial strain field and the mechanical property of the tissue. By using the IVUS strain field inside atherosclerotic plaques (De Korte et al. 2002; Doyley 2012; Maurice et al. 2004a, 2007), several elasticity reconstruction approaches have been developed to characterize the entire lesion (Baldeusung et al. 2007; Deléaval et al. 2013; Doyley 2012; Le Floch et al. 2009, 2010, 2012; Nayak et al. 2017; Poree et al. 2017; Richards and Doyley 2011; Richards et al. 2015; Tacheau et al. 2016) and to delineate a subendoluminal layer of the arterial wall called the palpography domain (Céspedes et al. 2000; Deléaval et al. 2013; Doyley et al. 2001; Schaar et al. 2005; Widynski et al. 2014). Both techniques aim to characterize the mechanical properties of atherosclerotic lesions and ultimately differentiate vulnerable from stable plaques.

In the latter elasticity-palpography technique originally developed by Céspedes et al. (2000) and Doyley et al. (2001), and revisited by our group to account for the anatomic shape of the VP (Deléaval et al. 2013), the apparent local stiffness of the subendoluminal arterial

wall layer was computed assuming that all plaque constituents are elastically isotropic. Even though these studies highlighted promising results, they failed to address the anisotropic mechanical behavior of the arterial wall and atherosclerotic lesion (Holzapfel et al. 2005).

Therefore, the purpose of this study was to extend the theoretical framework of the palpography technique implemented by Deléaval et al. (2013) to account for the anisotropic mechanical behavior of the arterial wall. Based on the continuum mechanics theory prescribing the radial strain field, a new anisotropic elasticity-palpography technique (AE-PT) was successfully applied to five modeled plaque geometries, ten coronary lesions and one healthy vessel of patients imaged *in vivo* with IVUS. An elastic anisotropy index (AI) was extracted.

## METHODS

Using images of patients who underwent coronary IVUS, cross-sectional geometries of ten atherosclerotic plaques and one healthy vessel were manually extracted based on the echogenicity aspect. Additionally, five idealized plaque morphologies were designed and simulated to validate the proposed AE-PT and to investigate the effects of cap thickness and palpography domain size on computed palpograms. Then, a 3-D finite-element (FE) analysis was conducted for each plaque geometry to compute the exact displacement and strain field distributions. To validate the robustness of palpograms, we synthesized radiofrequency (RF) echo frames by combining the computed FE displacement field with an ultrasound simulation model based on the Field II software (Jensen and Svendsen 1992). For all simulated IVUS RF sequences, the radial strain elastograms were estimated with the Lagrangian speckle model estimator (Maurice et al. 2004a). These radial strain elastograms were used to obtain the inverse solution of palpograms for the IVUS mechanical characterization of anisotropic vascular tissues.

### *IVUS study and plaque geometries*

*Patient population.* Vulnerable plaque geometries were extracted from IVUS images acquired on patients referred for percutaneous coronary intervention at the Louis Pradel Cardiology Hospital of Lyon after a first acute coronary syndrome with troponin I elevation. Patients' consent was obtained, and the study was approved by the ethical institutional review board of the Cardiology Hospital.

*Intravascular ultrasound imaging.* A set of one healthy and ten non-ruptured VP morphologies were collected from IVUS scans of the coronary arteries

according to the protocol described by Rioufol *et al.* (2002). Cross-sectional IVUS images of the atherosclerotic lesions were recorded at sites exhibiting the thinner fibroatheroma cap or at sites with the largest degree of stenosis. IVUS scans were performed with the iLab platform (Boston Scientific, Watertown, MA, USA) equipped with 40-MHz catheters (Atlantis SR Pro 3.6 F, Boston Scientific) and a radial spatial resolution in B-mode of approximately 90  $\mu\text{m}$  (Chopard *et al.* 2010).

*IVUS image analysis and plaque morphologic measurements.* IVUS echogenicity aspects were used to characterize VP components according to the following observations: (i) highly hypo-echogenic components (or anechogenic zones), suggestive of lipid or cellular deposition; or (ii) homogeneous reflective components, suggestive of fibrosis (Di Mario *et al.* 1998). A manual segmentation procedure using ImageJ software (National Institutes of Health [NIH], Bethesda, MD, USA) was performed by a cardiologist to extract the internal (*i.e.*, lumen) and external contours of the coronary cross section, as well as the contours of each plaque component (*i.e.*, necrotic core and dense fibrosis).

The morphology of each IVUS lesion image was quantified for plaque area ( $\text{Pla}_{\text{area}}$ ,  $\text{mm}^2$ ), lumen area ( $\text{Lu}_{\text{area}}$ ,  $\text{mm}^2$ ), soft necrotic core area, dense fibrosis area and degree of stenosis ( $\text{Stenos}_{\text{deg}}$ , %) as  $100 \times \text{Pla}_{\text{area}} / (\text{Pla}_{\text{area}} + \text{Lu}_{\text{area}})$  and cap thickness ( $\text{Cap}_{\text{thick}}$ , mm), which was defined as the shortest distance between the lumen and the necrotic core.

*Idealized geometries.* One concentric and one eccentric idealized circular plaque geometries with homogeneous anisotropic and quasi-incompressible atherosclerotic lesions were simulated to validate the theoretical background of the AE-PT (Fig. 1). Moreover, to investigate the effects of cap thickness ( $\text{Cap}_{\text{thick}}$ ) on the resulting anisotropic elasticity palpogram, plaque geometry acquired on a single patient (plaque #2 with a  $\text{Cap}_{\text{thick}}$  of 93  $\mu\text{m}$ ) (Fig. 2) was modified by translating the soft necrotic inclusion radially (*i.e.*, along the axis going from the center of mass of the vessel lumen to the center of mass of the necrotic core). Thus, three idealized plaque geometries with  $\text{Cap}_{\text{thick}}$  values of 200, 400 and 600  $\mu\text{m}$  were modeled (plaques #2A–2C) (Fig. 2).

*Forward problem: Spatial displacement and strain field distributions*

*3-D finite-element simulations.* The 2-D digitized contours obtained were transferred into Comsol software (Structural Mechanics Module, Version 4.3b, Comsol, France). To compute the deformed shape, a 3-D static FE simulation was conducted on each thin-sliced

volumetric plaque (slice thickness = 0.1 mm) obtained by extrusion of the cross-sectional plaque profile in the longitudinal direction of the artery. Each geometry was meshed with approximately 5000 six-node prism elements. The barycenter (*i.e.*, the center of mass) of the luminal area was used as the origin of the cylindrical coordinate system ( $R, \theta, Z$ ).

Instantaneous pressures were not recorded during the clinical IVUS exploration of the coronary arteries. Therefore, a blood pressure differential amplitude  $\Delta P$  between 0.3 and 1 kPa (*i.e.*, 2.2–7.5 mmHg) was assumed, corresponding to a realistic pressure gradient occurring between two successive images ( $\text{Im}^i$  and  $\text{Im}^{i+1}$ ) of the IVUS sequence (Le Floc'h *et al.* 2009, 2012). In our computational approach, the deformation of the atherosclerotic plaque between  $\text{Im}^i$  and  $\text{Im}^{i+1}$  was obtained by considering the geometry of the lesion at  $\text{Im}^i$  as the reference configuration. Therefore, FE simulations were performed in linear elasticity with a small loading blood pressure differential  $\Delta P$ . A free boundary condition was assumed at the external wall of the artery, and sliding conditions were imposed on the two cross sections of the vessel segment ( $R-\theta$  planes,  $Z=0$  and 0.1 mm). Such boundary conditions constrain the deformation to take place in the planes where  $Z$  is equal to a constant. Additionally, displacement limitations were placed upon nodes located on the four cardinal directions of the external wall of the artery to avoid rigid rotations. These latter boundary conditions allow free-radial expansion of the vessel during pressure loading, resulting in more realistic vessel inflation kinematics.

The fibrotic plaque medium and the healthy coronary arterial wall were considered as orthotropic and quasi-incompressible media (Appendix I), with initial mechanical properties coming from measurements (Chagnon *et al.* 2017; Holzapfel *et al.* 2005) of the pathologic coronary thick intima layer and the adventitia layer, respectively. The mechanical properties of the dense fibrosis and soft necrotic cores were considered as isotropic and quasi-incompressible (Lee *et al.* 1991; Loree *et al.* 1994). Young's moduli, Poisson ratios and shear moduli for all media are summarized in Table 1.

*Simulation of ultrasound images.* The open source software Field II (Jensen and Svendsen 1992) was modified to simulate the IVUS acoustic field (Fromageau *et al.* 2003; Nayak *et al.* 2017; Poree *et al.* 2017). Briefly, biological tissues were characterized by their acoustic scattering amplitude. Acoustic models were built by randomly distributing scatterers over each non-deformed medium. The relative acoustic scatterer amplitudes of blood, necrotic core, fibrotic plaque and dense fibrosis were set to 10%, 15%, 35% and 60%, respectively

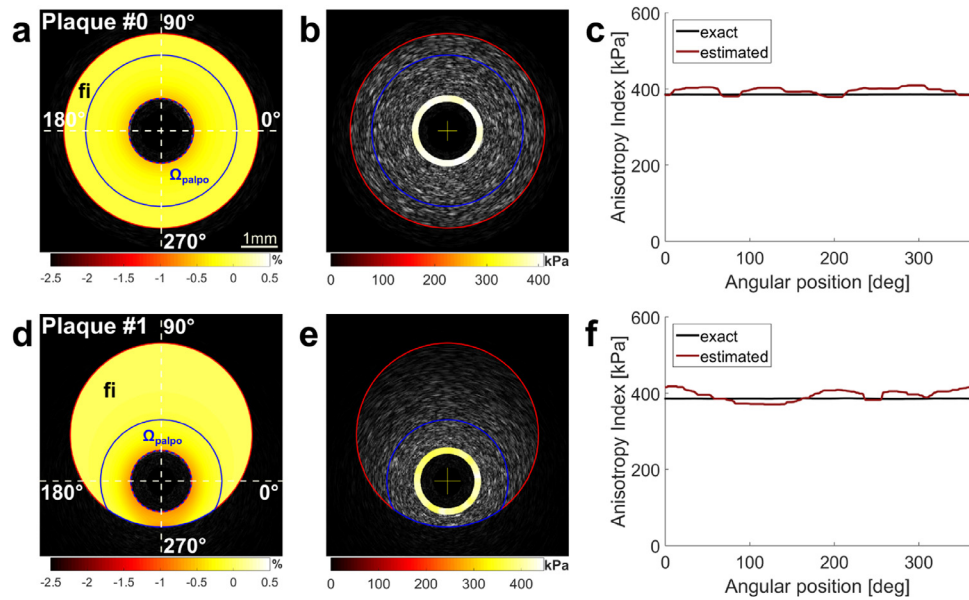


Fig. 1. Performance of the anisotropic elasticity-palpography technique on idealized homogeneous concentric and eccentric plaque models (plaques #0 and 1, respectively). (a, d) Plaque geometries (red contours), palpography domains  $\Omega_{\text{palpo}}$  (blue contours) and estimated radial strain elastograms. (b, e) Intravascular ultrasound images with anisotropy index-palpograms  $AI(\theta)$ . (c, f) Comparisons between exact and estimated  $AI(\theta)$  palpograms. fi = fibrotic plaque region.

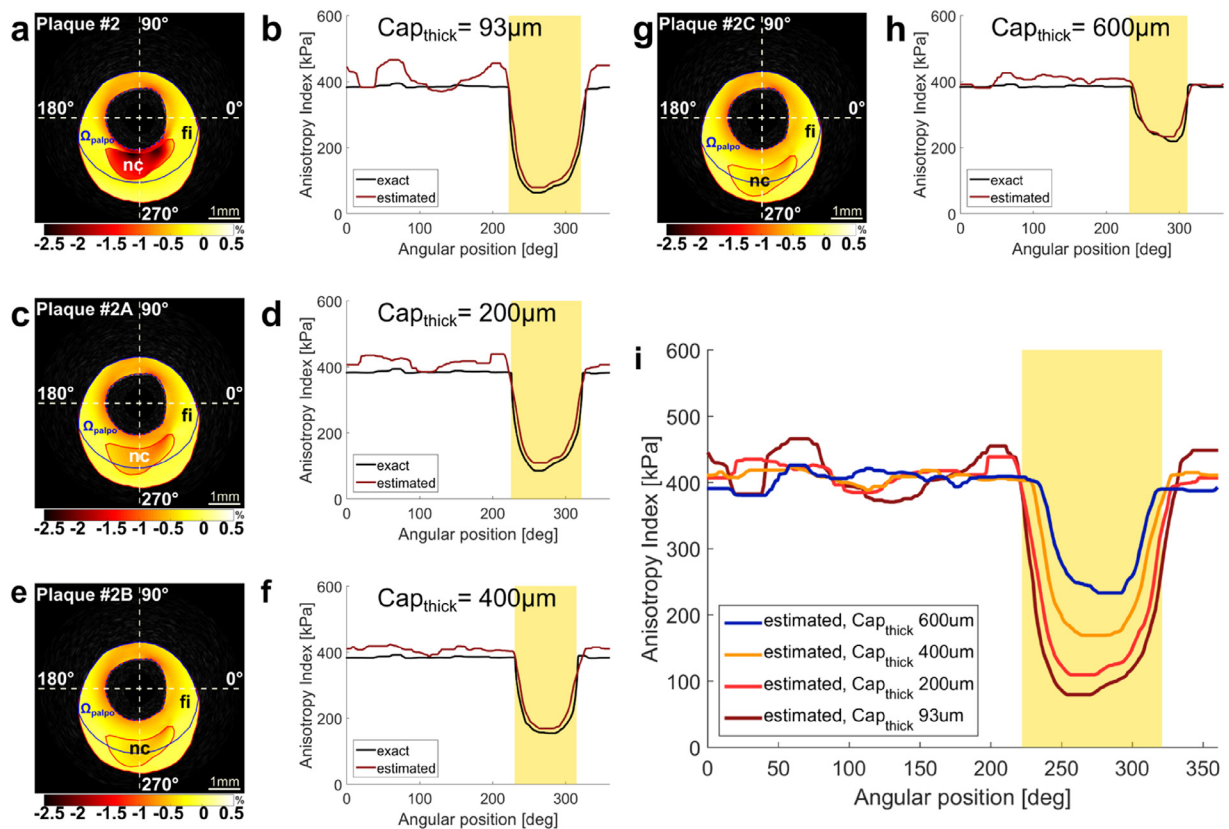


Fig. 2. Effect of cap thickness ( $Cap_{\text{thick}}$ ) on the resulting anisotropic elasticity palpogram  $AI(\theta)$ . Simulations were performed with plaque #2. (a, c, e, g) Plaque geometries (red contours), palpography domains (blue contours) and estimated radial strain elastograms. (b, d, f, h) Comparisons between exact and estimated  $AI(\theta)$  palpograms. (i) Comparison between all estimated  $AI$ -palpograms. Areas shaded in yellow indicate the location of the necrotic core. fi = fibrotic plaque region; nc = necrotic core.



Table 1. Orthotropic and isotropic mechanical properties and anisotropy index values assigned for the healthy wall, fibrous plaque, dense fibrosis and soft necrotic core inclusions

	Young's modulus (kPa)			Anisotropy index (kPa)	Reference (medium)
	$E_R$	$E_\theta$	$E_Z$		
Healthy wall	120	80	170	121.8	Holzapfel <i>et al.</i> (2005) (adventitia)
Pathologic wall					
Fibrous plaque	360	230	330	384.3	Holzapfel <i>et al.</i> (2005) (thick intima)
Dense fibrosis*		1500		2000	Lee <i>et al.</i> (1991)
Necrotic core*		5		6.7	Loree <i>et al.</i> (1994)

\* Isotropic ( $E_R = E_\theta = E_Z$ ).

(Cardoso *et al.* 2014; Le Floc'h *et al.* 2010; Maurice *et al.* 2004a). First, an IVUS image of the non-deformed state (*i.e.*,  $Im^i$ ) was created. Then the displacement field resulting from the FE simulation was used to track the location of scatterers and to create the post-deformation IVUS image (*i.e.*,  $Im^{i+1}$ ). A single rotating piezo-electric element (600- $\mu\text{m}$  diameter, 40-MHz central emission frequency) was modeled. An incremental circumferential rotation step of  $0.7^\circ$  was set to get 512 lines of RF signal for IVUS image reconstruction. A sampling frequency of 400 MHz was mimicked to represent the RF signal (SVMI HD IVUS system, ACIST Medical Systems, Eden Prairie, MN, USA). The arterial wall was also adjusted to set the centroid of the lumen coincident with the probe location.

*Intravascular strain elastography.* Sequences of simulated RF images were processed using the Lagrangian speckle model estimator. Radial strain fields were computed using a two-step strain imaging algorithm (Majdoulina *et al.* 2014; Maurice and Bertrand 1999; Maurice *et al.* 2004a, 2007). First, a block matching method provides local registration on overlapping sub-windows (measurement windows [MWs]). It allows compensating for rigid translation movements using a 2-D cross correlation coefficient. Once MW locations have been registered, the 2-D-deformation matrix is estimated. Estimation is based on the resolution of a first-order optical flow equation for each MW (Majdoulina *et al.* 2014). Sizes of MWs in radial and circumferential directions were set to 101 pixels (about 194.5  $\mu\text{m}$ ) and 21 pixels (about  $14.8^\circ$ ). Shift between MWs was set to preserve an overlap of 90% in the radial and circumferential directions (Le Floc'h *et al.* 2010; Maurice *et al.* 2004a). Moreover, a sensitivity study was conducted to investigate the influence of MW size on the accuracy of the AI-palpograms. In agreement with the IVUS imaging studies of Céspedes *et al.* (2000), Doyley *et al.* (2001), Deléaval *et al.* (2013), Widynsky *et al.* (2014) and Tacheau *et al.* (2016), a palpography domain thickness

close to 1 mm was used for our simulations, unless otherwise stated.

*Inverse problem: The anisotropic elasticity-palpography technique*

To account for the orthotropic mechanical properties of the vascular tissue, we defined an elastic anisotropy index (AI) as the ratio of the averaged difference between the radial and circumferential stresses ( $\sigma_{RR}$  and  $\sigma_{\theta\theta}$ ) over the averaged radial strain ( $\varepsilon_{RR}$ ) along the radial axis:

$$AI(\theta) = \frac{\left| \int_{R_i(\theta)}^{R_o(\theta)} \{ \sigma_{RR}(R, \theta) - \sigma_{\theta\theta}(R, \theta) \} dR \right|}{\varepsilon(\theta)} \quad (1)$$

with

$$\varepsilon(\theta) = \left| \int_{R_i(\theta)}^{R_o(\theta)} \varepsilon_{RR}(R, \theta) dR \right| \quad (2)$$

where  $R_i(\theta)$  and  $R_o(\theta)$  are the inner and outer radii of the considered palpography domain, respectively. Appendix II provides the details of the computation of this index.

Note that the amplitude of  $AI(\theta)$  remains constant and equal to

$$AI(\theta) = E_R^0 \left[ 1 - \frac{1}{4} \left( 1 - \frac{E_R^0}{E_\theta^0} + \frac{E_R^0}{E_Z^0} \right) \left( 1 + \frac{E_Z^0}{E_R^0} - \frac{E_Z^0}{E_\theta^0} \right) \right]^{-1} \quad (3)$$

when considering a homogeneous pathologic lesion made of the same incompressible orthotropic medium (*i.e.*, when  $E_R(R, \theta) = E_R^0$ ,  $E_\theta(R, \theta) = E_\theta^0$  and  $E_Z(R, \theta) = E_Z^0$ ), whatever the plaque geometry and the palpography domain  $\Omega_{\text{palpo}}$  considered (with  $R_i(\theta) \leq R \leq R_o(\theta)$ ).

For a homogeneous incompressible transversely isotropic lesion (*i.e.*, when  $E_R(R, \theta) = E_R^0$  and  $E_Z(R, \theta) = E_\theta(R, \theta) = E_\theta^0$ ), the AI amplitude is reduced to the constant

$$AI = \frac{4E_R^0{}^2}{4E_R^0 - E_\theta^0} \quad (4)$$

and to  $AI = 4E/3$  for a homogeneous incompressible isotropic plaque (*i.e.*, when  $E_R(R, \theta) = E_\theta(R, \theta) = E_Z(R, \theta) = E$ ).

Table 1 gives the amplitudes of the AI for the considered mechanical properties of the adventitia (healthy arterial wall), thick intima (fibrotic plaque), dense fibrosis and soft necrotic media.

*Method used to verify the accuracy of the anisotropic elasticity-palpography technique.* Two anisotropic index palpograms were computed: the first,  $AI(\theta)$ , with the estimated radial strain elastogram, and the second,  $AI^*(\theta)$ , considering the exact radial strain map. To investigate the accuracy of each palpogram (*i.e.*,  $AI(\theta)$  and  $AI^*(\theta)$ ), we compared them with the exact circumferential distribution of the averaged anisotropic material constant  $C(R, \theta)$  along the radial axis:

$$AI^{\text{exact}}(\theta) = \frac{1}{R_o(\theta) - R_i(\theta)} \int_{R_i(\theta)}^{R_o(\theta)} C(R, \theta) dR \quad (5)$$

and we computed two mean relative anisotropy index errors ( $MR_{\text{error}}^{AI}$  and  $MR_{\text{error}}^{AI^*}$ ):

$$MR_{\text{error}}^{K(\theta)} (\%) = \frac{100}{2\pi} \int_{\theta=0}^{2\pi} \left| \frac{K(\theta) - AI^{\text{exact}}(\theta)}{AI^{\text{exact}}(\theta)} \right| d\theta, \quad (6)$$

with  $K(\theta) = AI(\theta)$  or  $AI^*(\theta)$

$MR_{\text{error}}^{AI^*}$  indicates the best accuracy that could be reached when using the new anisotropic palpography technique, and  $MR_{\text{error}}^{AI}$  quantifies the cumulative error induced by both the proposed palpography technique and the estimated radial strain elastogram. Therefore, to quantify the error induced only by the radial strain elastogram estimation on the AI-palpogram, we defined the following mean relative error ( $MR_{\text{error}}^{AI/AI^*}$ ):

$$MR_{\text{error}}^{AI/AI^*} (\%) = \frac{100}{2\pi} \int_{\theta=0}^{2\pi} \left| \frac{AI(\theta) - AI^*(\theta)}{AI^*(\theta)} \right| d\theta \quad (7)$$

## RESULTS

### IVUS study

Seven non-ruptured VPs with soft necrotic cores, one homogeneous plaque, two atherosclerotic lesions with dense fibrosis inclusions and one non-pathologic cross section were identified after extensive IVUS scanning. The geometric characteristics of these 11 cross sections scanned *in vivo* are summarized in Table 2. All computed AI-palpograms presented in the figures were obtained from radial strain elastograms estimated from sequences of simulated RF images.

### Performance of the anisotropic elasticity-palpography technique

#### Validation cases and AI-palpogram accuracies.

Simulations were performed on the idealized concentric and eccentric plaque geometries with homogeneous orthotropic and quasi-incompressible fibrotic atherosclerotic plaques (Fig. 1). AI-palpograms for the concentric and eccentric plaques remain almost constant with mean values ( $\pm$ standard deviation [SD]) of  $396 \pm 9$  and  $394 \pm 14$  kPa, both close to the expected theoretical value ( $AI = 384.3$  kPa, Table 1) given by eqn (3)—even in the high-strained free-plaque arc length of the eccentric plaque geometry, which is the thinnest region of the palpography domain (*i.e.*, between the angular positions  $180^\circ$  and  $360^\circ$ , Figs. 1e, f). The small amplitudes of the mean relative anisotropy index errors ( $MR_{\text{error}}^{AI} = 3.08\%$  and  $3.70\%$ , Table 3), together with the small amplitudes of the standard deviations ( $SD = 1.62\%$  and  $2.08\%$ , Table 3), highlight the robustness of the AE-PT algorithm with respect to the IVUS RF sequences when estimating AI-palpograms for concentric and eccentric plaque morphologies, respectively. More generally, the results revealed that all 18 AI-palpograms were estimated with reasonable accuracy (with a mean relative error  $MR_{\text{error}}^{AI} = 26.79 \pm 48.84\%$ , Table 3) when compared with the exact ones and with a good robustness with respect to the estimated strain-elastogram, as reflected by the mean relative error  $MR_{\text{error}}^{AI/AI^*} = 10.24 \pm 14.43\%$  (Table 3).

*Detection and characterization of vulnerable plaques with soft inclusions.* AI-palpograms were computed for VPs with one (plaques #2, 2A–2C and 12–15, Figs. 2–4) and two distinct (plaques #16 and 17, Fig. 5) necrotic cores. These AI-palpograms were estimated with a mean relative error of  $34.36 \pm 57.03\%$  (Table 3). Locations of the necrotic core sites were accurately detected using the AE-PT. The AI-palpogram amplitudes were found close to the AI value of the fibrotic plaque (*i.e.*, close to 384.3 kPa, Table 1) except at soft inclusion locations, for which the AI amplitudes were found to be significantly lower, as expected (Figs. 2–4).

*Detection and characterization of stable plaques with dense fibrosis inclusions.* For one stable plaque (plaque #19, Fig. 6) with one isolated dense fibrosis inclusion, the proposed palpography technique detected and distinguished the fibrotic plaque from dense fibrosis. Compared with the exact value, the estimated AI amplitude at the dense fibrosis inclusion location was underestimated by almost a factor of 2.

Table 2. Characteristics of the IVUS-detected and modelled atherosclerotic plaques and healthy artery used to test the performance of the proposed anisotropic elasticity–palpography technique.

Plaque No.	Origin of geometry	Cap thickness* ( $\mu\text{m}$ )	Area ( $\text{mm}^2$ )			Lumen	Stenosis (%)
			Necrotic core	Dense fibrosis	Plaque		
0	Model	—	—	—	14.14	1.77	89
1	Model	—	—	—	25.13	3.14	89
2	IVUS	93	1.36	—	8.74	2.79	76
2A	Model	200	1.36	—	8.74	2.79	76
2B	Model	400	1.36	—	8.74	2.79	76
2C	Model	600	1.36	—	8.74	2.79	76
Healthy 1	IVUS	—	—	—	8.45	11.87	42
11	IVUS	—	—	—	9.88	7.59	57
12	IVUS	<90 (55)	3.48	—	14.24	6.06	70
13	IVUS	<90 (63)	4.41	—	15.48	4.28	78
14	IVUS	<90 (49)	1.97	—	19.38	6.80	74
15	IVUS	<90 (48)	1.89	—	17.29	8.44	67
16	IVUS	100	1.41/0.46	—	19.50	7.67	72
17	IVUS	<90 (65)	2.20/0.53	—	29.53	4.34	87
18	IVUS	<90 (60)	—	1.36	6.34	5.99	51
19	IVUS	<90 (88)	—	0.90	5.31	5.35	50

IVUS = intravascular ultrasound.

\* Cap thickness was randomly assigned (value in parentheses) when found to be under the limit of the B-mode spatial resolution obtained with the 40-MHz IVUS catheter (*i.e.*, <90  $\mu\text{m}$ ).

*Detection and characterization of homogeneous stable plaques and healthy vessel wall.* AI-palpograms were also computed for one eccentric homogeneous stable plaque (plaque #11, Fig. 7) and one healthy coronary wall (healthy #1, Fig. 8). The AE-PT successfully characterized the orthotropic mechanical properties of the pathologic lesion (made of fibrotic plaque tissue) and of the non-pathologic coronary artery. The two AI-palpograms were found almost constant with mean values ( $\pm\text{SD}$ ) of  $360 \pm 15$  and  $134 \pm 9$  kPa, close to the

expected theoretical values: AI = 384.3 and 121.8 kPa for the pathologic and healthy arteries, respectively (Table 1).

#### *Effects of Cap<sub>thick</sub> on estimated AI-palpogram.*

For a given palpography domain, the Cap<sub>thick</sub> significantly affects the AI-palpogram at the soft inclusion location (plaques #2 and 2A–2C, Fig. 2). Simulations performed with the idealized VP models in which the Cap<sub>thick</sub> increases from 93 to 600  $\mu\text{m}$  revealed that the

Table 3. Accuracy of the proposed anisotropic elasticity-palpography technique and effect of the estimated radial strain elastogram (computed with a measurement window size of  $101 \times 21$  pixels) on the estimated AI-palpograms\*

Plaque No.	NC	DF	H	MR <sub>error</sub> <sup>AI</sup> (%)	MR <sub>error</sub> <sup>AI</sup> (%)	MR <sub>error</sub> <sup>AI/AI*</sup> (%)
0	—	—	✓	0.22 $\pm$ 0.14	3.08 $\pm$ 1.62	2.9 $\pm$ 1.49
1	—	—	✓	3.72 $\pm$ 1.72	3.7 $\pm$ 2.08	4.24 $\pm$ 3.2
2	✓	—	—	4.86 $\pm$ 4.53	13.41 $\pm$ 9.45	8.76 $\pm$ 6.06
2A	✓	—	—	5.97 $\pm$ 5.26	11.13 $\pm$ 7.9	5.53 $\pm$ 4.28
2B	✓	—	—	3.74 $\pm$ 2.29	7.84 $\pm$ 4.35	4.22 $\pm$ 3.06
2C	✓	—	—	1.91 $\pm$ 1.55	5.05 $\pm$ 3.85	4.03 $\pm$ 2.79
Healthy 1	—	—	✓	0.65 $\pm$ 0.5	10.03 $\pm$ 7.67	10.49 $\pm$ 7.66
11	—	—	✓	2.45 $\pm$ 2.24	6.66 $\pm$ 3.55	5.91 $\pm$ 3.43
12	✓	—	—	21.71 $\pm$ 27.64	33.16 $\pm$ 52.64	16.4 $\pm$ 19.1
13 (0.75)	✓	—	—	66.6 $\pm$ 52.63	83.23 $\pm$ 71.73	16.1 $\pm$ 21.47
13	✓	—	—	59.43 $\pm$ 57.53	74.14 $\pm$ 67.96	15.98 $\pm$ 18.61
13 (1.25)	✓	—	—	36.96 $\pm$ 41.17	46.45 $\pm$ 55.68	15.84 $\pm$ 17.44
14	✓	—	—	35.21 $\pm$ 62.45	44.87 $\pm$ 78.06	9.96 $\pm$ 11.05
15	✓	—	—	35.54 $\pm$ 73.87	45.19 $\pm$ 92.07	8.49 $\pm$ 7.47
16	✓	—	—	8.72 $\pm$ 9.57	14.46 $\pm$ 17.38	8.11 $\pm$ 9.05
17	✓	—	—	24.86 $\pm$ 22	33.3 $\pm$ 33.19	8.36 $\pm$ 9.24
18	—	✓	—	11.32 $\pm$ 12.96	35.34 $\pm$ 31.27	29.5 $\pm$ 31.75
19	—	✓	—	3.76 $\pm$ 6.38	11.25 $\pm$ 8.91	9.43 $\pm$ 6.69
Mean $\pm$ SD				18.2 $\pm$ 37.99	26.79 $\pm$ 48.84	10.24 $\pm$ 14.43

AI = anisotropy index; DF = dense fibrosis; H = homogeneous; NC = necrotic core; SD = standard deviation.

\* See eqns (6) and (7) for definitions of the mean relative errors MR<sub>error</sub><sup>AI</sup>, MR<sub>error</sub><sup>AI</sup> and MR<sub>error</sub><sup>AI/AI\*</sup>. Palpography thickness is 1 mm unless otherwise indicated in parentheses beside the plaque number.

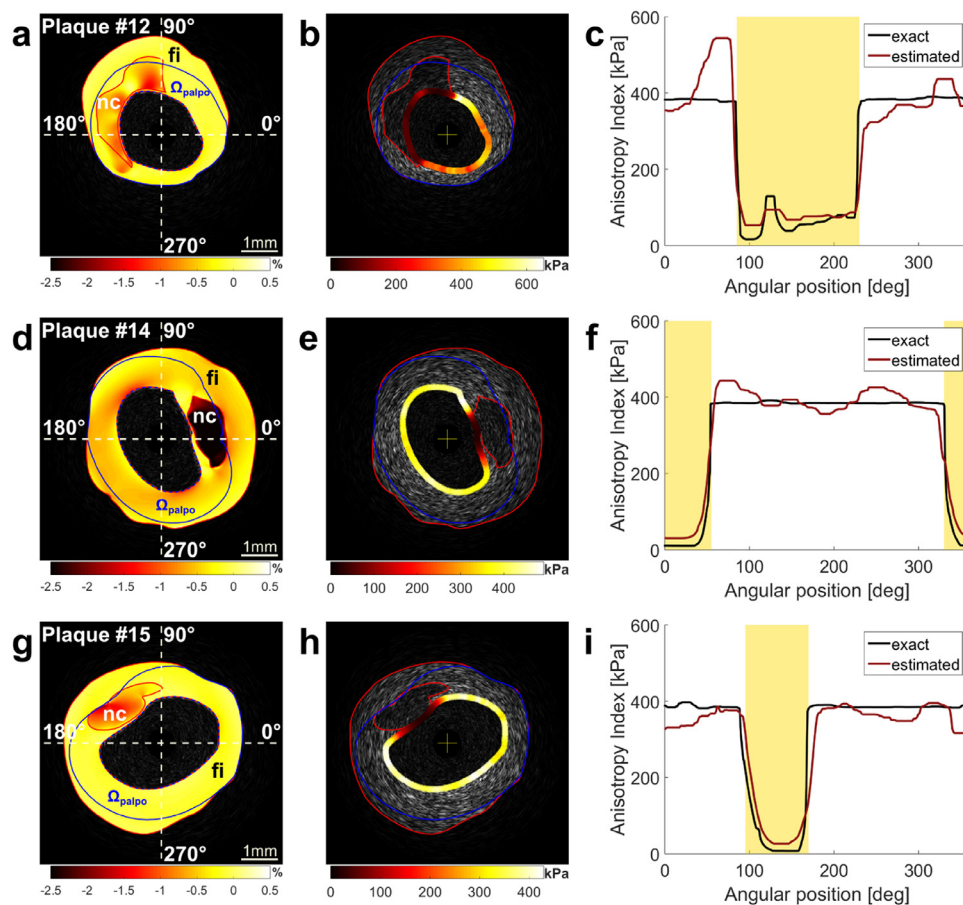


Fig. 3. Performance of the anisotropic elasticity palpography technique in detection of vulnerable plaques with one necrotic core (plaques #12, 14 and 15). (a, d, g) Plaque geometries (*red contours*), palpography domains (*blue contours*) and estimated radial strain-elasticograms. (b, e, h) IVUS images with anisotropy index-palpograms  $AI(\theta)$ . (c, f, i) Comparisons between exact and estimated  $AI(\theta)$  palpograms. Areas shaded in *yellow* indicate the locations of the necrotic cores. fi = fibrotic plaque region; nc = necrotic core.

mean AI amplitude at the location of the necrotic core increased from  $139 \pm 74$  to  $300 \pm 65$  kPa (Fig. 2).

*Effects of the palpography domain size on estimated AI amplitude.* We computed the AI-palpograms for the same VP considering three palpography domains with maximum thicknesses of 0.75, 1 and 1.25 mm (plaque #13, Fig. 4). Increasing the palpography domain thickness from 0.75 to 1.25 mm improved the accuracy of the AI palpogram from  $MR_{\text{error}}^{\text{AI}} = 83.23 \pm 71.73\%$  to  $MR_{\text{error}}^{\text{AI}} = 46.45 \pm 55.68\%$ , respectively. The location of the necrotic core site was accurately detected using all palpography domains.

*Sensitivity of AI-palpograms to the accuracy of the estimated strain elastograms.* Because the MW size is one of the main parameters driving the accuracy of the computed strain elastograms, we restricted our sensitivity analysis to this parameter. These

computations were performed on the stable plaque #18 with one dense fibrosis inclusion and by considering three MW sizes:  $75 \times 15$  pixels (*i.e.*,  $144.4 \mu\text{m} \times 10.6^\circ$ ),  $101 \times 21$  pixels (*i.e.*,  $194.5 \mu\text{m} \times 14.8^\circ$ ) and  $121 \times 31$  pixels (*i.e.*,  $232.9 \mu\text{m} \times 21.8^\circ$ ). An important palpogram variation was found at the free-plaque arc length. Only for this particular plaque, the AI-palpogram obtained with  $(101 \times 21)$  pixels MW size fail to characterize the atherosclerotic lesion, indicating that the accuracy of the AI-palpogram depends on the MW size used when estimating the radial strain elastogram (Fig. 9).

## DISCUSSION

The new intraluminal ultrasonic palpation imaging technique successfully quantified the apparent wall stiffness of the anisotropic vascular wall based on the arterial radial strain and blood pressure data set.



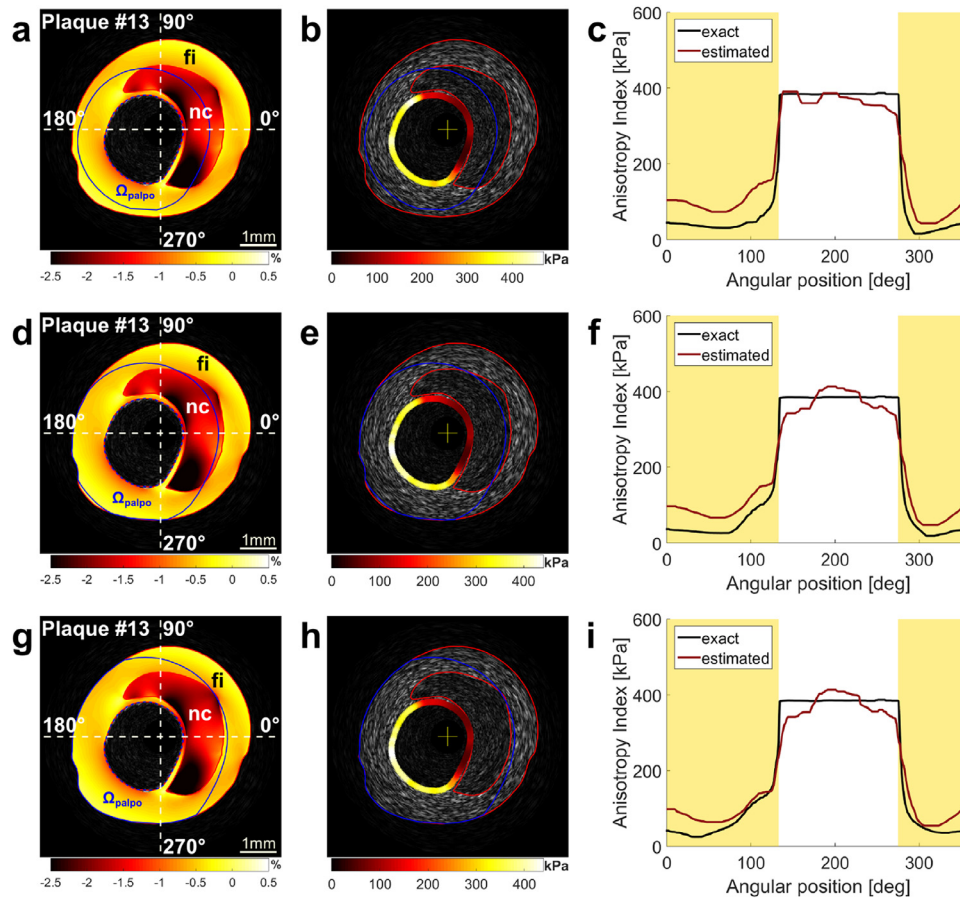


Fig. 4. Effect of palpography domain size on the resulting anisotropic elasticity palpogram. Simulations were performed with plaque #13 considering three palpography domain thicknesses. (a, d, g) Plaque geometries (*red contours*); estimated radial strain-elasticograms and palpography domains with thicknesses of 0.75, 1 and 1.25 mm, respectively (*blue contours*). (b, e, h) Intravascular ultrasound images with anisotropy index-palpograms  $AI(\theta)$ . (c, f, i) Comparisons between exact and estimated  $AI(\theta)$  palpograms. Areas shaded in *yellow* indicate the location of the necrotic core. fi = fibrotic plaque region; nc = necrotic core.

#### Physical meaning of the elastic anisotropy index $AI$

One of the advantages of AI, which may be viewed as an anisotropic apparent modulus, is to account for the three Young's moduli of the orthotropic atherosclerotic lesion. Even though the AE-PT does not allow identifying individually the three orthotropic material constants, this index, which combines the three orthotropic Young's moduli, is proportional to the radial Young's modulus and depends on the two stiffness ratios:  $k_1 = E_\theta / E_R$  and  $k_2 = E_Z / E_R$ . Equation (3) indicates that the normalized elastic anisotropy index  $AI/E_R$  increases with  $k_2$  and decreases with  $k_1$  when  $0.5 \leq k_i \leq 2$  ( $i = 1, 2$ ). In other words, an increase in AI may be due to either an increase in  $E_R$  and  $E_Z$  or a decrease in  $E_\theta$ . Furthermore, the AI amplitude appears to be very sensitive to the presence of mechanical heterogeneities, dropping significantly when predicting soft inclusions in the composite atherosclerotic lesions (Figs. 2–5).

#### AE-PT highlights the non-linear mechanical response of the vascular tissue

Because the proposed constitutive model refers to the configuration of the plaque at  $Im^i$ , rather than the initial zero-stress configuration, the linear tangent elasticity procedure used in our approach (Bonet and Wood 1997) remains acceptable as long as the absolute amplitudes of the incremental strain components estimated between the two successive IVUS images  $Im^i$  and  $Im^{i+1}$  remain small ( $\leq 2\%$ ) (Le Floch *et al.* 2012). Therefore, the computed AI must be viewed as an anisotropic linearized tangent modulus (Holzapfel 2000) measured for the physiologic pressure range considered. The other advantage of such an incremental linearized technique applied on a series of IVUS images is that it allows quantification of the non-linear mechanical response of the fibrotic plaque tissue during the physiologic increase in blood pressure.

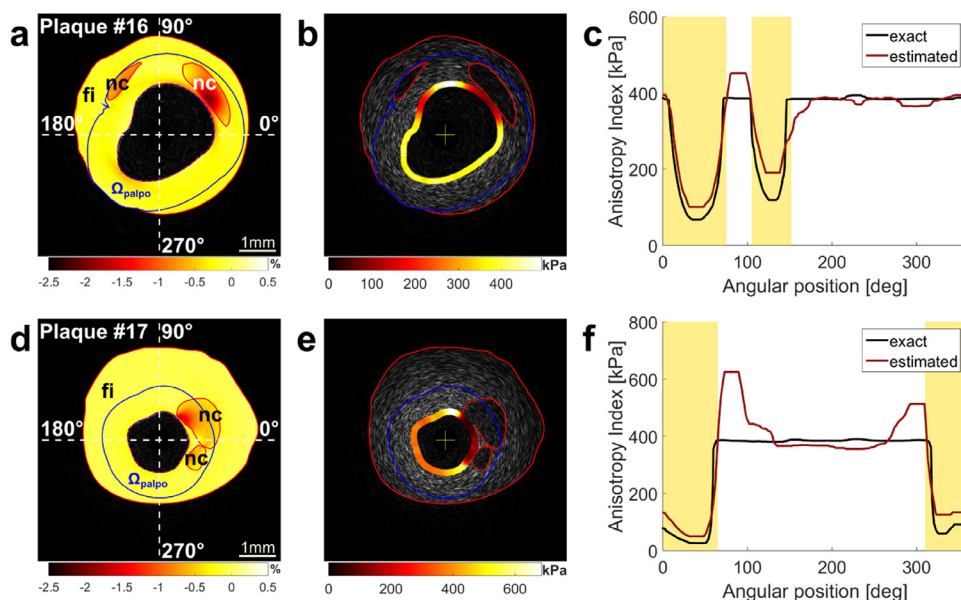


Fig. 5. Performance of the anisotropic elasticity-palpography technique in detection of vulnerable plaques with two necrotic cores (plaques #16 and 17). (a, d) Plaque geometries (*red contours*), palpography domains (*blue contours*) and estimated radial strain elastograms. (b, e) intravascular ultrasound images with anisotropy index-palpogram  $AI(\theta)$ . (c, f) Comparisons between exact and estimated  $AI(\theta)$  palpograms. Areas shaded in *yellow* indicate the locations of the two necrotic cores. fi = fibrotic plaque region; nc = necrotic core.

#### How should cardiologists choose the palpography domain?

With the AE-PT, the interventional cardiologist may define the palpography domain by considering any arbitrary endoluminal thick layer size and shape. The results clearly indicated that an increase in the palpography domain thickness does not necessarily decrease the accuracy of the AI-palpogram (see Table 3 and Fig. 4, plaque #13). This could be explained by the fact that the palpography domain must be thin enough to obtain a reasonable signal-to-noise (SNR) ratio (Céspedes et al. 2000; Doyley et al. 2001) and large enough for a good performance of the strain imaging algorithm (Maurice et al. 2004a).

#### What is the advantage of using the AI instead of the fractional anisotropy index for characterizing a VP?

The principal strain and fractional strain anisotropy indexes are two other strain-based metrics that were defined and applied in non-invasive clinical cardiovascular magnetic resonance imaging and ultrasound studies (Nayak et al. 2017; Soleimanifard et al. 2010). However, extracting the anisotropic mechanical properties of the tissue from these indexes is difficult. Furthermore, although the estimations of these indexes require accurate knowledge of all strain tensor components, estimation of the AI requires only the radial strain component, which is the only component that can be extracted from the coronary RF-IVUS signal with reasonable accuracy.

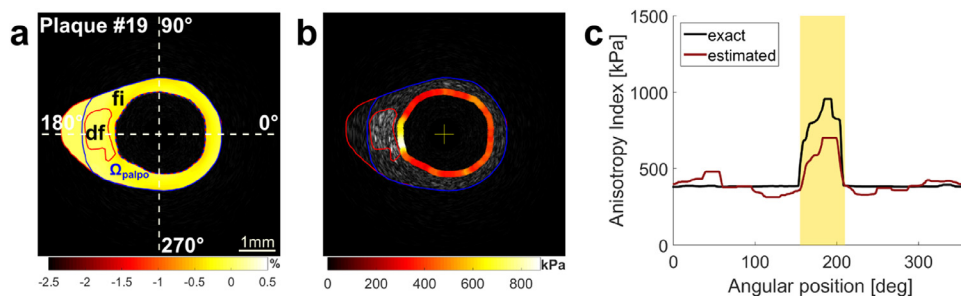


Fig. 6. Performance of the anisotropic elasticity-palpography technique in detection of a stable plaque with one dense fibrosis inclusion (plaque #19). (a) Plaque geometry (*red contour*), palpography domain (*blue contour*) and estimated radial strain elastogram. (b) Intravascular ultrasound images with anisotropy index-palpogram  $AI(\theta)$ . (c) Comparison between exact and estimated  $AI(\theta)$  palpograms. Area shaded in *yellow* indicates the location of the dense fibrosis inclusion. df = dense fibrosis. fi = fibrotic plaque region.

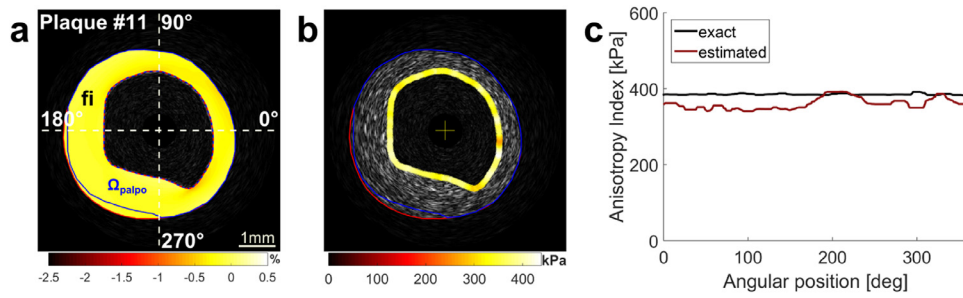


Fig. 7. Performance of the anisotropic elasticity-palpography technique in detection of a stable homogeneous fibrotic plaque (plaque #11). (a) Plaque geometry (*red contours*), palpography domain (*blue contours*) and estimated radial strain elastogram. (b) Intravascular ultrasound image with the anisotropy index-palpogram  $AI(\theta)$ . (c) Comparison between the exact and estimated  $AI(\theta)$  palpograms. *fi* = fibrotic plaque region.

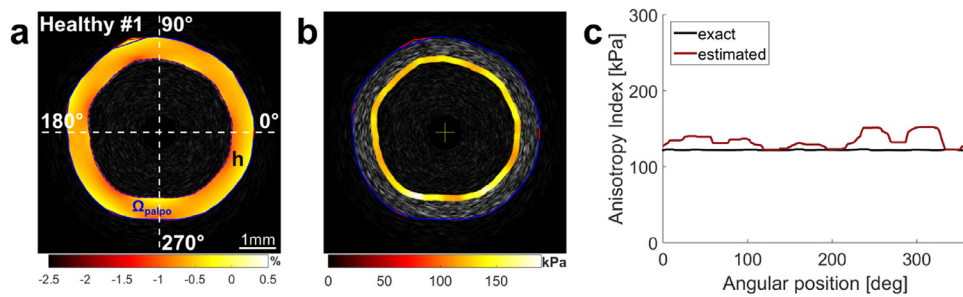


Fig. 8. Performance of the anisotropic elasticity-palpography technique in characterization of a healthy coronary cross section (healthy 1). (a) Artery geometry (*red contours*), palpography domain (*blue contours*) and estimated radial strain elastogram. (b) Intravascular ultrasound image with the anisotropy index-palpogram  $AI(\theta)$ . (c) Comparison between the exact and the estimated  $AI(\theta)$  palpograms.

*Is the AE-PT sufficient to detect VPs and diagnose plaque rupture?*

The proposed AE-PT cannot diagnose the degree of plaque vulnerability to rupture because it does not allow

measurements of other key geometric factors (Cilla *et al.* 2012; Ohayon *et al.* 2008). However, this technique remains reliable for detecting and highlighting complex VPs with one or several soft inclusions (see Figs. 2–5).

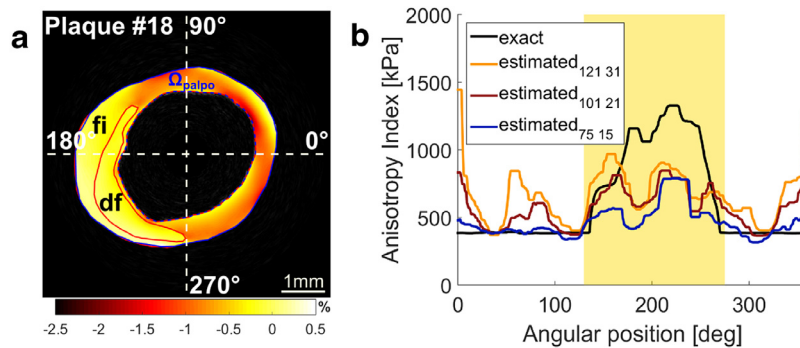


Fig. 9. Sensitivity analysis revealing the influence of measurement window (MW) size on the computed anisotropy index-palpogram  $AI(\theta)$ . All these computations were performed on the dense fibrotic plaque #18. Three MW sizes were considered for these simulations:  $75 \times 15$  pixels,  $101 \times 21$  pixels and  $121 \times 31$  pixels. (a) Artery geometry (*red contour*), palpography domain (*blue contour*) and estimated radial strain map obtained with the  $101 \times 21$ -pixels MW size. (b) Comparison between exact and estimated  $AI(\theta)$  palpograms. The area shaded in *yellow* indicates the location of the dense fibrosis inclusion. *df* = dense fibrosis; *fi* = fibrotic plaque region.

### Study limitations

Several limitations deserve to be pointed out. First, the anisotropic palpation technique was tested and validated only on synthesized RF echo frames. Although further animal and clinical studies are needed to extend and strengthen the present findings, our preliminary results appear to be promising and indicate the potential and feasibility of this novel imaging technique. The second limitation of this study is that the palpography model neglected the residual strain, which has a significant effect on the stress amplitude in the thin fibrous cap (Ohayon et al. 2007). However, because the AI is an anisotropic linearized tangent modulus (Holzapfel 2000) measured during physiologic conditions, it implicitly accounts for all existing internal stresses/strains, including the residual ones. Finally, one clinical case presented in this study (plaque #18, Fig. 9) highlighted the importance of considering optimal MW sizes to compute accurate AI-palpograms. A more exhaustive study clearly deserves to be investigated as future work to define an MW size that would optimize the performance of the strain-imaging algorithm of Majdoulina et al. (2014).

### Relevance to other clinical applications

Modifying the mechanical properties of the vulnerable atherosclerotic lesion could be the key issue for the stabilization of VPs. Finet et al. (2004) reported that a very slight increase in soft necrotic core stiffness can tilt a VP from instability to stability. Today, therapies based on statins (Libby et al. 2002; Nozue et al. 2012) and other emergent medications (Nicholls et al. 2016) are used by clinicians to enhance plaque stability. However, the effects of all these drugs on the mechanical properties of coronary atherosclerosis lesions still remain unclear. The proposed improved anisotropic elasticity-palpography imaging technique could be used by clinicians to investigate the evolution of the anisotropic mechanical properties of atherosclerotic plaques during drug therapies.

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### CONFLICT OF INTEREST DISCLOSURE

The authors have no conflicts of interest to disclose.

## APPENDIX I. CONSTITUTIVE LAW IN LINEAR ELASTICITY FOR INCOMPRESSIBLE ORTHOTROPIC VASCULAR TISSUES

The constitutive law is given using the cylindrical coordinate system  $(R, \theta, Z)$ . In linear elasticity, the relationships between the material constants of a compressible orthotropic medium and the stress tensor  $(\sigma_{ij}$  with  $i, j = R, \theta, Z)$  and strain tensor  $(\varepsilon_{ij}$  with  $i, j = R, \theta, Z)$  components are given by Berthelot (1999):

$$\begin{Bmatrix} \varepsilon_{RR} \\ \varepsilon_{\theta\theta} \\ \varepsilon_{ZZ} \\ 2\varepsilon_{\theta Z} \\ 2\varepsilon_{RZ} \\ 2\varepsilon_{R\theta} \end{Bmatrix} = \begin{bmatrix} 1/E_R & -\nu_{\theta R}/E_R & -\nu_{ZR}/E_Z & 0 & 0 & 0 \\ -\nu_{R\theta}/E_R & 1/E_\theta & -\nu_{Z\theta}/E_Z & 0 & 0 & 0 \\ -\nu_{RZ}/E_R & -\nu_{\theta Z}/E_\theta & 1/E_Z & 0 & 0 & 0 \\ 0 & 0 & 0 & 1/G_{\theta Z} & 0 & 0 \\ 0 & 0 & 0 & 0 & 1/G_{RZ} & 0 \\ 0 & 0 & 0 & 0 & 0 & 1/G_{R\theta} \end{bmatrix} \begin{Bmatrix} \sigma_{RR} \\ \sigma_{\theta\theta} \\ \sigma_{ZZ} \\ \sigma_{\theta Z} \\ \sigma_{RZ} \\ \sigma_{R\theta} \end{Bmatrix} \quad (\text{AI.1})$$

where  $E_i$  ( $i = R, \theta, Z$ ) are the three Young's moduli associated with the  $i$  directions,  $G_{ij} = G_{ji}$  ( $i, j = R, \theta, Z$  with  $i \neq j$ ) are the three shear moduli associated with the  $ij$  planes, and  $\nu_{ij} = -\varepsilon_{ij}/\varepsilon_{ii}$  ( $i, j = R, \theta, Z$  with  $i \neq j$ ) are the six Poisson's ratios (i.e. the ratios of the strain in the  $j$  direction to the strain in the perpendicular  $i$  direction). Such orthotropic material also obeys to the reciprocity conditions:  $\nu_{ij}/E_i = \nu_{ji}/E_j$  ( $i, j = R, \theta, Z$  with  $i \neq j$ ). Thanks to these relationships, the number of material constants drops to nine. Moreover, assuming that the vascular wall incompressibility imposes the following three relationships among the Poisson's ratios:

$$\nu_{R\theta} + \nu_{RZ} = 0, \quad \nu_{\theta R} + \nu_{\theta Z} = 0 \quad \text{and} \quad \nu_{ZR} + \nu_{Z\theta} = 0. \quad (\text{AI.2a} - 2c)$$

So knowing the three Young's moduli  $E_R$ ,  $E_\theta$  and  $E_Z$ , we can extract the six Poisson's ratios.

## APPENDIX II. THE ELASTIC ANISOTROPY INDEX AI

Assuming the vascular media as incompressible and under plane strain constraint ( $\varepsilon_{RZ} = \varepsilon_{\theta Z} = \varepsilon_{ZZ} = 0$ ), one can derive the following relationships (Berthelot 1999):

$$\sigma_{RR}(R, \theta) - \sigma_{\theta\theta}(R, \theta) = C(R, \theta)\varepsilon_{RR}(R, \theta) \quad (\text{AII.1a})$$

with

$$C(R, \theta) = E_R(R, \theta)[1 - \nu_{RZ}(R, \theta)\nu_{ZR}(R, \theta)]^{-1} \quad (\text{AII.1b})$$

$$\nu_{RZ}(R, \theta) = \frac{1}{2} \left[ 1 - \frac{1}{k_1} + \frac{1}{k_2} \right] \quad (\text{AII.1c})$$

$$\nu_{ZR}(R, \theta) = \frac{1}{2} \left[ 1 + k_2 - \frac{k_2}{k_1} \right] \quad (\text{AII.1d})$$



$$k_1 = \frac{E_\theta(R, \theta)}{E_R(R, \theta)} \quad (\text{AII.1e})$$

and

$$k_2 = \frac{E_Z(R, \theta)}{E_R(R, \theta)} \quad (\text{AII.1f})$$

where,  $\sigma_{RR}(R, \theta)$ ,  $\sigma_{\theta\theta}(R, \theta)$  and  $\varepsilon_{RR}(R, \theta)$  are the radial and circumferential stress and radial strain components, respectively.

In this linearized elasticity approach, we assumed that the incremental radial strain  $\varepsilon_{RR}(R, \theta)$  is proportional to the imposed change in pressure  $\Delta P$  and inversely proportional to the anisotropic material constant  $C(R, \theta)$ :

$$\varepsilon_{RR}(R, \theta) = \frac{\Delta P}{C(R, \theta)} g(R, \theta) \quad (\text{AII.2})$$

where  $g(R, \theta)$  is a correcting shape function, which account for the entire plaque morphology including regions of plaque heterogeneities. By substituting Eq. (AII.2) into Eq. (AII.1a), Eq. (AII.1a) can be rewritten as follow:

$$\sigma_{RR}(R, \theta) - \sigma_{\theta\theta}(R, \theta) = \Delta P g(R, \theta) \quad (\text{AII.3})$$

Taking advantage of this expression (Eq. (AII.3)), the elastic anisotropy index AI (Eq. (1)) becomes

$$AI(\theta) = \frac{\Delta P \left| \int_{R_i(\theta)}^{R_o(\theta)} g(R, \theta) dR \right|}{\varepsilon(\theta)} \quad (\text{AII.4})$$

Since we assumed that the unknown morphological function  $g(R, \theta)$  does not depend on the mechanical properties and because this function escapes to direct measurements, we approximated it by a correcting *homogeneous* shape function  $g^*(R, \theta)$  accounting only for the anatomical cross-sectional plaque geometry. This approximated correcting function was obtained by performing a FE simulation (with a loading blood pressure differential  $\Delta P$ ) in which the pathologic vessel was assumed homogeneous and made of a quasi-incompressible isotropic material. This approximated shape function  $g^*(R, \theta)$  was used to estimate the anisotropic palpography index:

$$AI(\theta) = \frac{\Delta P \left| \int_{R_i(\theta)}^{R_o(\theta)} g^*(R, \theta) dR \right|}{\varepsilon(\theta)} \quad (\text{AII.5})$$

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