Vector flow imaging of the ascending aorta. Are systolic backflow and atherosclerosis related?

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Vector flow imaging of the ascending aorta

Are systolic backflow and atherosclerosis related?

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Abstract—In the ascending aorta, atherosclerotic plaque formation, which is a risk factor for cerebrovascular events, most often occurs along the inner curvature. Atherosclerosis is a multifactorial disease, but the predilection site for the aortic vessel degradation is probably flow dependent. To better understand the aortic flow and especially the complex flow patterns, the ascending aorta was scanned intraoperatively in patients undergoing heart surgery using the angle-independent vector velocity ultrasound method Transverse Oscillation (TO). The primary aim of the study was to analyze systolic backflow in relation to atherosclerosis. Thirteen patients with normal aortic valves were included in to the study. TO implemented on a conventional US scanner (ProFocus 2202 UltraView, BK Medical, Herlev, Denmark) with a linear array transducer (8670, BK Medical, Herlev, Denmark) was used intraoperatively on the ascending aorta in long axis view. The presence of systolic backflow, visualized with TO, was correlated to aortic atherosclerosis, to systolic velocities obtained with transesophageal echocardiography and cardiac output obtained with pulmonary artery catheter thermodilution, to gender, age, aortic diameter, left ventricular ejection fraction (LVEF) and previous myocardial infarctions (MI). Systolic backflow in the ascending aorta was present for 38% (n=5) of the patients. The location of the backflow was strongly associated to the location of the plaques (p<0.005), and backflow was associated to high systolic velocities (p<0.05). The other obtained parameters were not associated to systolic backflow. It was shown that systolic backflow is a common flow feature in the ascending aorta, and that backflow is associated to atherosclerotic plaques and systolic velocities. The study indicates that vector flow imaging using TO can provide important blood flow information in the assessment of atherosclerosis.

Keywords—vector velocity imaging; Transverse Oscillation; ascending aorta; systolic backflow; atherosclerosis

I. INTRODUCTION

Atherosclerosis is a disease of the vessel wall with accumulation of low-density lipo-proteins and migration of macrophages, resulting in plaques formation and wall thickening. Despite numerous advances in medical and surgical vascular interventions, atherosclerosis still is the leading cause of death in the Western world [1]. The development of the disease is multi-factorial, however, the predilection sites for plaques formation are probably mainly governed by vessel geometry and blood flow [2].

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II. MATERIALS AND METHODS

A. Transverse Oscillation

The TO method estimates both the axial and transverse velocity component with emission of a conventional pulse for Doppler ultrasound. The motion in the axial direction is found as in conventional Doppler ultrasound, while the motion in the transverse direction is found by using a changed apodization in receive beamforming with a special estimator (Fig. 1) [13]. Previous papers provide detailed explanations of the method [5, 6, 13].

A conventional ultrasound scanner (ProFocus 2202 UltraView, BK Medical, Herlev, Denmark) and a linear transducer (8670, BK Medical, Herlev, Denmark) under sterile settings were used to record epiaortic scan sequences in long axis view (Fig. 2).

The angle-independent vector velocities are displayed in real-time on the B-mode image as colored pixels given by a 2-D color bar defining both direction and velocity magnitude. Small arrows superimposed on to the color flow map are added to facilitate flow interpretation (Fig. 3).

The temporal resolution of TO estimation is 16 frames/s, and the maximum scan depth is approximately 5 cm due to the transducer setup available. For each scan session, the color box was adjusted to cover the lumen, and the pulse repetition frequency, depth setting, gain, and wall filtering were adjusted for vector velocity estimation. Each scan sequence corresponded to 15 s.

B. Transesophageal echocardiography and pulmonary artery catheter thermodilution

Transesophageal echocardiography (TEE) was performed on a Philips iE33 scanner equipped with a Philips X7-2t transducer (Philips Healthcare, Eindhoven, The Netherlands) with the scan parameters adjusted for flow imaging. The scan sequences were obtained through the aortic valve aperture in the deep transgastric long axis view with the line of interrogation placed along the left ventricular outlet tract and the ascending aorta.

Pulmonary artery catheter thermodilution (PACTD) was performed with a 7.5F pulmonary artery catheter (Swan-Ganz Oximetry TD catheter, Baxter Edwards Critical Care, Irvine, CA, USA) and the cardiac output was determined by the HP-Philips M1012A Cardiac Output Module (Hewlett Packard, Boeblingen, Germany) as a mean of four bolus injections of 10 ml saline. TO and PACTD measurements were performed simultaneously, while TO and TEE measurements were performed within 5 min. For each patient, the medical record was consulted to obtain information on previous MI. LVEF was estimated prior to the cardiac surgery with conventional echocardiography.

The presence and placement of systolic backflow were analyzed by visual inspection of the vector map, and the presence and placement of atherosclerotic plaques were noted by inspecting the B-mode sequence. The diameter of the ascending aorta was obtained by measuring the wall-to-wall distance in short axis view.

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C. Statistics

Data were analyzed with descriptive statistics and comparisons were calculated with one-way ANOVA and Fisher’s exact test depending on variable types. A p-value of less than 0.05 was considered significant. Statistical analyses were performed with IBM SPSS Statistics v. 19 (SPSS Inc., Chicago, IL, USA).

III. RESULTS

The study was approved by the local ethics committee (no. H-2-2012-039). Thirteen patients (12 males and 1 female, mean age 66.1 y.o., range 41-80 y.o.) without any history of valvular disease undergoing coronary bypass surgery entered the study after written informed consent.

TABLE I. DESCRIPTIVE STATISTICS

| Backflow | Number of patients | Mean age [years] | Gender | PACTD [mm] | Diameter [mm] | LVEF | Age | MI | PACTD | TEE | LVEF | Plaques | Gender | Age | P | Prior |
|----------|-------------------|------------------|--------|------------|---------------|------|-----|----|--------|-----|------|---------|--------|-----|---|---|------|
| Present  | 5                 | 70.6 (8.2)       | 4/1    | 4.0 (1.4)  | 134.0 (39.1)  | 32.6 (3.1) | 4/1 |
| Not present | 8           | 63.3 (12.1)      | 8/0    | 4.9 (1.2)  | 101.3 (14.3)  | 31.8 (5.8) | 1/7 |
| Total    | 13                | 66.1 (11.0)      | 12/1   | 4.5 (1.3)  | 113.8 (30.1)  | 32.1 (4.8) | 5/8 |

* The patients are grouped according to the presence of systolic backflow. Mean or frequency (with the standard deviation) is given for each variable.

Systolic backflow in the ascending aorta was present for 38% (n=5) of the patients (Table 1) and observed during the entire systole. Of the patients with backflow, three patients had backflow along the inner convexity of the aortic arch, one along the outer concavity and one along both (Table 2). Backflow was significantly associated to high systolic velocities measured by TEE (p<0.05) but not to PACTD (p<0.52) (Table 3). The location of the plaques was associated to the location of the backflow as illustrated in Fig. 1 (p<0.005) and shown in Table 2 and 3. All performed comparisons are given in Table 3, and as shown, no other significant associations were found.

TABLE II. SYSTOLIC BACKFLOW AND PLAQUE LOCATION

<table>
<thead>
<tr>
<th>Atherosclerotic plaque location</th>
<th>None</th>
<th>Concave vessel wall</th>
<th>Convex vessel wall</th>
<th>Both vessel walls</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Systolic backflow location</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>None</td>
<td>7</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>8</td>
</tr>
<tr>
<td>Concave vessel wall</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Convex vessel wall</td>
<td>0</td>
<td>0</td>
<td>3</td>
<td>0</td>
<td>3</td>
</tr>
<tr>
<td>Both vessel walls</td>
<td>0</td>
<td>1</td>
<td>0</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>Total</td>
<td>8</td>
<td>1</td>
<td>4</td>
<td>0</td>
<td>13</td>
</tr>
</tbody>
</table>

Systolic backflow was seen for 38% of the patients, and the backflow was observed during the entire systole as reported previously [11, 12]. There was a strong association between the location of the backflow and location of atherosclerotic plaques (Table 2 and 3). Systolic backflow and atherosclerotic plaques were primarily found along the inner convex curvature (Table 2 and Fig. 3) as reported by others [2]. Moreover, high systolic velocities were related to the presence of backflow (Table 3).

The complex patterns in the blood flow are recognized as an essential factor for atherogenesis, and the blood flow-induced wall shear stress is thought to play a major role, as regions with low wall shear stress are predilection sites for atherosclerosis [14]. The site of plaque formation in the thoracic aorta has in previous studies been linked to retrograde flow patterns [15], and Svedlund et al. showed increased occurrence of aortic arch atheromas in patients with pronounced diastolic retrograde flow [16]. In this study, a strong association was found between the location of systolic backflow, presumably with low wall shear stress, and the location of plaques formation, and between the presence of backflow and high systolic velocities.

Bensalah et al. found systolic backflow associated to age and aortic diameter [17], however it was not found in this study, but might be a consequence of a relatively small sample size, as age seems to differ slightly between groups (Table 1).

No patient was classified with aortic stenosis, however, narrowing of the left ventricular outlet tract (LVOT) probably created the jet with high systolic velocities and adjacent systolic backflow. Thus, narrowing of LVOT may be one of the risk factors for atherogenesis in the ascending aorta, through the formation of systolic backflow as previously reported [18]. Further studies are warranted to explore this hypothesis.

The intraoperative scan approach with the Transverse Oscillation system was chosen to overcome the limited penetration depth. Hence, efforts are made for implementation of TO on phased and curved array transducers for increased penetration [19, 20]. It is evident that blood flow in the ascending aorta is highly complex and non-axisymmetrical.

IV. DISCUSSION
Therefore, using 2-D vector flow only provides a partial visualization of the aortic blood flow, and to fully understand the aortic flow, 3-D US is warranted. 3-D vector flow using TO has been obtained, but is not yet implemented for clinical evaluation [21, 22].

With the introduction of TO in to cardiac US imaging, new insonation windows of the heart will be possible due to the angle-independency, and new information of the complex flow patterns of the cardiovascular system can be obtained.

Ischemic strokes have been linked to atheromas in the thoracic aorta [23] and the presence of retrograde flow has likewise been suspected to increase the risk of retrograde embolization to the cerebrum [16]. Therefore, evaluation of complex flow patterns in the ascending aorta is important, and the angle-independent flow estimation provided by TO is a promising visualization modality for this purpose.

Only one radiologist (KLH) evaluated the B-mode and TO sequences, which is a potential confounder. Furthermore, the relatively small study population limits this study. To confirm the presented preliminary results, a study including a larger study population with healthy aortic valve is on-going, in which, the blood flow of the ascending aorta is examined in both short and long axis view with TO.

V. CONCLUSION

The complex flow in the ascending aorta among 13 patients was investigated in real-time and intraoperatively with the vector flow method TO. Systolic backflow was seen in 38% of the patients, and the backflow was present during the entire systole. The systolic backflow was strongly associated with the location of atherosclerotic plaques, moderately associated to peak systolic velocity, but not to cardiac output, LVEF, previous MI, gender, age or aortic diameter.

Thus, the study indicates that systolic backflow and atherosclerotic plaque formation in the ascending aorta are related. The evaluation of systolic backflow may predict, which patients are prone to atherosclerotic plaque formation in the ascending aorta, and thereby, be a tool for prevention of ischemic stroke. Hence, a better understanding of the complex flow patterns in the ascending aorta may be achieved, and flow-related mechanisms in relation to aortic diseases such as atherosclerosis can be explored with the angle-independent vector velocity method TO.

References