ORIGINAL ARTICLE

Haemodynamic and anatomic progression of aortic stenosis

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ABSTRACT

Aortic valve stenosis (AS) is a progressive disease, but the impact of baseline AS haemodynamic or anatomic severity on AS progression remains unclear.

Methods In 149 patients (104 mild AS, 36 moderate AS and 9 severe AS) enrolled in 2 ongoing prospective cohorts (COFRASA/GENERAC), we evaluated AS haemodynamic severity at baseline and yearly, thereafter, using echocardiography (mean pressure gradient (MPG)) and AS anatomic severity using CT (degree of aortic valve calcification (AVC)).

Results After a mean follow-up of 2.9±1.0 years, mean MPG increased from 22±11 to 30±16 mm Hg (+8±3 mm Hg/year), and mean AVC from 1108±891 to 1640±1251 AU (arbitrary units) (+188±176 AU/year). Progression of AS was strongly related to baseline haemodynamic severity (+2±3 mm Hg/year in mild AS, +4±3 mm Hg/year in moderate AS and +5±5 mm Hg/year in severe AS (p=0.01)), and baseline haemodynamic severity was an independent predictor of haemodynamic progression (p=0.0003). Annualised haemodynamic and anatomic progression rates were significantly correlated (r=0.55, p<0.0001), but AVC progression rate was also significantly associated with baseline haemodynamic severity (+14±133 AU/year in mild AS, +279±189 AU/year in moderate AS and +361±293 AU/year in severe AS, p<0.0001), and both baseline MPG and baseline AVC were independent determinants of AVC progression (p<0.0001).

Conclusions AS progressed faster with increasing haemodynamic or anatomic severity. Our results suggest that a medical strategy aimed at preventing AVC progression may be useful in all subsets of patients with AS including those with severe AS and support the recommended closer follow-up of patients with AS as AS severity increases.

Clinical trial registration COFRASA (clinicalTrials.gov number NCT00338676) and GENERAC (clinicalTrials.gov number NCT00647088).

INTRODUCTION

Aortic valve stenosis (AS) is the most common valvular heart disease in Western countries.1 AS affects 2%–7% of people aged over 70 years2,3 and its prevalence is going to dramatically increase with the ageing of the population. AS haemodynamic severity is usually assessed using echocardiography based on mean gradient, peak velocity (PV) and calculation of the aortic valve area (AVA).4 Aortic valve calcification (AVC) is the main process leading to AS, and can be accurately and quantitatively measured (calcium score) in vivo using multislice CT (MSCT).5,6 The degree of AVC—AS anatomic severity—is closely related to AS haemodynamic severity as assessed using echocardiography and can be considered as a complementary method for the evaluation of severity of AS in difficult clinical situations.7,8

Calcific AS is a progressive disease,9 and despite recent progress, determinants of AS progression remain unclear. AS progression is highly variable among individuals, and the impact of baseline—haemodynamic or anatomic—AS severity remains unknown. Thus, in 2 ongoing prospective cohorts, we aimed to evaluate the impact of baseline AS severity assessed, either by using echocardiography (haemodynamic assessment) or MSCT (anatomic assessment) on AS progression.

METHODS

Study design

Patients with degenerative AS, enrolled between November 2006 and May 2013 in two ongoing prospective studies, COFRASA (clinicalTrials.gov number NCT00338676) and GENERAC (clinicalTrials.gov number NCT00647088), with at least 2 years of follow-up were considered in the present study. COFRASA and GENERAC aim at evaluating the determinants of AS occurrence and progression, and all participants underwent a comprehensive clinical, transthoracic echocardiographic (TTE) and MSCT evaluation at study entry and yearly thereafter. Inclusion criteria are pure, at least mild (defined by a mean pressure gradient (MPG) ≥10 mm Hg and aortic valve structural changes (thickening/calcification) asymptomatic AS. Exclusion criteria were AS due to rheumatic disease or radiotherapy, previous infective endocarditis, more than mild coexisting aortic regurgitation (defined by a vena contracta width ≥3 mm or a regurgitant volume ≥30 mL) or associated valvular disease and severe renal insufficiency (creatinine clearance ≤30 mL/min). Echocardiographic and MSCT measurements were performed with each blinded from the other. The study was approved by our regional ethics committee, and all patients gave written informed consent.

Echocardiography

A comprehensive Doppler echocardiogram was performed at baseline and then on a yearly basis.
Severity of AS was evaluated based on PV MPG and the A VA calculated using the continuity equation⁴ ¹¹ as recommended by current guidelines. The A VA was calculated as an absolute value and indexed (AVAi) to body surface area (BSA). Mild AS was defined by an MPG <25 mm Hg, moderate AS was defined by an MPG between 25 and 40 mm Hg and severe AS by an MPG >40 mm Hg. Short-axis view in systole was used to differentiate bicuspid from trileaflet aortic valve.

MSCT measurements
MSCT was performed on the same day as TTE at baseline and also yearly thereafter using a Philips scanner (MX 8000 IDT 16, Philips Medical Systems, Andover, Massachusetts, USA) or a General Electric scanner (Light speed VCTTM, General Electric Company, Fairfield, Connecticut, USA). A scan run consisted of a prospective acquisition of forty 3 mm thick contiguous transverse slices. MSCT was performed with a tube current of 75A and a voltage of 120 kV. Acquisition time was 0.5 s/slice ECG triggered at 75% of the RR interval. No contrast enhancement and a voltage of 120 kV . Acquisition time was 0.5 s/slice ECG triggered at 75% of the RR interval. No contrast enhancement was needed nor was a β-blocker administered for the purpose of the examination. Measurements were performed using dedicated semiautomatic software (Heart Beat Calcium Scoring, Philips Medical Systems or SmartScore, General Electric Medical Systems). Calcification was defined as four adjacent pixels with density greater than 130 Hounsfield units. The degree of A VC was quantitatively assessed according to the Agatston method (calcium score) and expressed in arbitrary units (AU). A VC was defined as calcification within the valve leaflets including the aortic annulus and the aortic wall immediately adjacent to the leaflets. Two MSCT runs were performed sequentially with a 1 mm or 2 mm initial interval. Each run was independently scored and the two scores were averaged. Radiation exposure was typically between 2 mSv and 3 mSv.

The accuracy and reproducibility of MSCT assessment of the degree of calcification has been previously validated. Intraobserver variability was 4±10% (68±130 AU) and interobserver variability was 6±7% (74±77 AU). Scans with motion artefacts precluding A VC measurements were excluded.

Statistical analysis
Continuous variables were expressed as mean±SD, median (25th and 75th percentile), or number of patients (per cent). Correlations between calcium score and AS haemodynamic severity were tested with linear and non-linear regressions, and the model using the square root of A VC was retained as providing the best fit. Annualised progression was calculated as (final measurement—baseline measurement)/follow-up duration for haemodynamic (MPG) and anatomic (A VC score) measurements. Comparisons between terciles of A VC progression were performed using Kruskal–Wallis test, Mann–Whitney–Wilcoxon test, χ² test or Fisher exact test, as appropriate. Linear regressions in univariate analysis and in multivariate analysis after adjustment for age, gender and aortic valve anatomy (bicuspid or trileaflet aortic valve) were used to determine the association between anatomic or haemodynamic progression (as a continuous variable) and baseline haemodynamic severity or baseline A VC score. A p value <0.05 was considered statistically significant.

RESULTS
Characteristics of the population
One hundred and forty-nine patients with at least mild AS and at least 2 years of follow-up were prospectively enrolled. Baseline characteristics of the population are summarised in table 1 (left part). Briefly, mean age was 74±9 years, 75% were male and 93% in sinus rhythm. The aortic valve was trileaflet in 125 patients (84%) and bicuspid in 24 (16%). MPG was 22±11 mm Hg, but there was a wide range of AS severity (median 20 mm Hg, (10–55)) and 104 patients had mild AS (MPG <25 mm Hg), 36 patients moderate AS (MPG between 25 to 40 mm Hg) and 9 patients severe AS (MPG >40 mm Hg). Mean A VC score was 1108±891 AU (median 873, (81–3923)). As previously reported, there was a good correlation between the degree of A VC assessed using MSCT and haemodynamic parameters of AS severity assessed either using the MPG (r=0.67, p<0.0001; figure 1), the PV (r=0.68; p<0.0001) or the A VAi (r=0.47; p<0.0001).

Haemodynamic progression
Mean follow-up was 2.9±1.0 years. At the end of follow-up, mean MPG increased to 30±16 mm Hg (median 25, (10–75)) and the yearly mean MPG increase was 3±3 mm Hg/year (median 2, (0–13)). Twelve patients progressed from mild AS to severe AS and 27 patients from moderate AS to severe AS. However, AS progression rate was not uniform and was strongly related to baseline haemodynamic severity (MPG, PV or A VAi at baseline, all r >0.29; p<0.0001). Thus, AS progression rate was 2±3 mm Hg/year (median 2, (0–13)) in patients with mild AS, 4±3 mm Hg/year (median 3, (0–15)) in patients with moderate AS and 5±5 mm Hg/year (median 4, (0–14)) in patients with severe AS (p=0.01) (figure 2A). By contrast, progression was not different between patients with trileaflet and bicuspid aortic valve (3±3 mm Hg/year (median 2, (0–13)) vs 3±4 mm Hg/year (median 2, (0–13)), p=0.70) or between male and female patients (3±3 mm Hg/year (median 2, (0–13)) vs 4±4 mm Hg/ year (median 3, (0–15)), p=0.40). After adjustment for age, gender and aortic valve anatomy (bileaflet or trileaflet aortic valve), baseline haemodynamic severity was an independent predictor of haemodynamic progression (p=0.0003). The same results were obtained when PV or A VAi were used instead of MPG.

Anatomic progression
Final mean A VC score was 1640±1251 AU (median 1341, (196–5246)), and mean A VC increase was 188±176 AU/year (median 141, (0–787)). There was no difference in A VC progression between patients with trileaflet and bicuspid aortic
valve (179±167 AU/year (median 135, (0–752)) vs 240 ±212 AU/year (median 201, (0–809)), p=0.30) or between male and female patients (196±166 AU/year (median 152, (0–650)) vs 163±201 AU/year (median 107, (0–820)), p=0.06). AVC progression rate was also significantly associated with baseline haemodynamic severity, and AVC progression was 141 ±133 AU/year (median 100, (0–582)) in patients with mild AS, 279±189 AU/year (median 241, (0–827)) in patients with moderate AS and 361±293 AU/year (median 230, (70–809)) in patients with severe AS (p<0.0001) (figure 2B). There was a correct linear correlation between anatomic and haemodynamic changes (r=0.55; p<0.0001; figure 3). However, as for haemodynamic progression, AVC progression was strongly related to baseline AVC load (r=0.57; p<0.0001; figure 4).

We then divided our population into terciles of AVC progression (<85 AU/year, 85–205 AU/year and >205 AU/year). Characteristics of the 149 patients according to AVC terciles of progression are presented in table 1 (right part). Baseline haemodynamic severity (MPG, PV and AVAi) and AVC score increased with AVC terciles (all p<0.001). Mean MPG increase was also different between terciles (1±1 mm Hg in the first tercile, 3±2 mm Hg in the second tercile and 5±4 mm Hg in the third tercile, p<0.0001). Haemodynamic severity and AVC at baseline were significant predictors of AVC progression in univariate analysis (both p<0.0001), whereas gender (p=0.33), age (p=0.69) and valve anatomy (p=0.20) were not. After adjustment for age, gender and valve anatomy, baseline AVC was an independent determinant of AVC progression (p<0.0001).

**DISCUSSION**

In this prospective cohort of patients with AS who were followed for up to 5 years, we observed that neither haemodynamic nor anatomic progression were uniform. There was a positive linear correlation between haemodynamic and anatomic progression, but both haemodynamic and anatomic severity at baseline were independent predictors of AS progression, whereas gender and aortic valve anatomy (bicuspid or trileaflet aortic valve) were not.

**AVC progression**

If more data accumulate showing that AS is an active process, AS progression remains unpredictable. Similarties with atherosclerosis and bone remodelling have been reported,13–16 but statin
therapy has failed to prevent AS progression in three clinical trials.17–19 Annualised AS haemodynamic progression is classically considered as constant and homogeneous (+7 mm Hg/year and +3 mm Hg/year in the Simvastatin and Ezetimibe in Aortic Stenosis (SEAS) trial20) even if it is highly variable among individuals. However, determinants of AS progression are largely unknown, and the impact of AS severity has rarely been evaluated. We have previously shown in a population-based study that the degree of AVC was an important predictor of AVC progression, but the sample size was relatively small and the impact of baseline haemodynamic severity could not be assessed.20 In the present study, we show a strong association between AS haemodynamic or anatomical progression and either baseline AVC score or baseline MPG, PV or AVA and baseline anatomical or haemodynamic severity was an independent predictor of AS progression. Thus, baseline AS severity was a major determinant of AS dynamic severity was an independent predictor of AS progression.

Baseline characteristics of the overall population overall and according to terciles of progression of aortic valve calcification (AVC)

<table>
<thead>
<tr>
<th>Terciles of AVC progression</th>
<th>Overall n=149</th>
<th>First tercile (&lt;85 AU/year)</th>
<th>Second tercile (85–205 AU/year)</th>
<th>Third tercile (&gt;206 AU/year)</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, years</td>
<td>74±9</td>
<td>73±9</td>
<td>75±8</td>
<td>75±10</td>
<td>0.2</td>
</tr>
<tr>
<td>Body surface area, m²</td>
<td>1.9±0.2</td>
<td>1.9±0.2</td>
<td>1.8±0.2</td>
<td>1.9±0.2</td>
<td>0.1</td>
</tr>
<tr>
<td>Male gender</td>
<td>112 (75)</td>
<td>35 (70)</td>
<td>35 (71)</td>
<td>42 (84)</td>
<td>0.2</td>
</tr>
<tr>
<td>Sinus rhythm</td>
<td>138 (93)</td>
<td>48 (96)</td>
<td>45 (92)</td>
<td>45 (90)</td>
<td>0.5</td>
</tr>
<tr>
<td>Diabetes</td>
<td>32 (21)</td>
<td>9 (18)</td>
<td>14 (29)</td>
<td>9 (18)</td>
<td>0.34</td>
</tr>
<tr>
<td>Hypertension</td>
<td>102 (69)</td>
<td>32 (64)</td>
<td>33 (67)</td>
<td>37 (74)</td>
<td>0.55</td>
</tr>
<tr>
<td>Smoker</td>
<td>80 (54)</td>
<td>26 (52)</td>
<td>27 (55)</td>
<td>27 (54)</td>
<td>0.95</td>
</tr>
<tr>
<td>Trileaflet aortic valve</td>
<td>125 (84)</td>
<td>43 (86)</td>
<td>43 (88)</td>
<td>39 (78)</td>
<td>0.4</td>
</tr>
<tr>
<td>Total cholesterol</td>
<td>4.76±1.10</td>
<td>4.78±1.21</td>
<td>4.60±0.86</td>
<td>4.90±1.09</td>
<td>0.36</td>
</tr>
<tr>
<td>Low-density lipoprotein cholesterol</td>
<td>2.67±0.94</td>
<td>2.73±0.97</td>
<td>2.43±0.86</td>
<td>2.84±0.95</td>
<td>0.06</td>
</tr>
<tr>
<td>Serine creatinine (μmol/L)</td>
<td>92±29</td>
<td>97±36</td>
<td>87±21</td>
<td>92±28</td>
<td>0.67</td>
</tr>
<tr>
<td>Calcium (mmol/L)</td>
<td>2.37±0.12</td>
<td>2.37±0.13</td>
<td>2.38±0.12</td>
<td>2.36±0.11</td>
<td>0.35</td>
</tr>
<tr>
<td>Phosphate (mmol/L)</td>
<td>1.06±0.17</td>
<td>1.08±0.18</td>
<td>1.08±0.15</td>
<td>1.02±0.18</td>
<td>0.26</td>
</tr>
<tr>
<td>EF, %</td>
<td>63±5</td>
<td>63±6</td>
<td>63±6</td>
<td>63±4</td>
<td>0.5</td>
</tr>
<tr>
<td>Baseline AVC score, cm²</td>
<td>1.41±0.36</td>
<td>1.57±0.38</td>
<td>1.38±0.36</td>
<td>1.29±0.30</td>
<td>0.0007</td>
</tr>
<tr>
<td>Baseline indexed AVC area, cm²/m²</td>
<td>0.75±0.18</td>
<td>0.82±0.19</td>
<td>0.75±0.19</td>
<td>0.68±0.15</td>
<td>0.0007</td>
</tr>
<tr>
<td>Baseline MPG, mm Hg</td>
<td>22±11</td>
<td>17±9</td>
<td>20±6</td>
<td>29±13</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Baseline peak aortic velocity, cm/s</td>
<td>299±62</td>
<td>270±55</td>
<td>290±44</td>
<td>338±68</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Baseline AVC score, AU</td>
<td>1108±891</td>
<td>775±761</td>
<td>825±459</td>
<td>171±1020</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Mean MPG increase, mm Hg/year</td>
<td>3±3</td>
<td>1±1</td>
<td>3±2</td>
<td>5±4</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Mean AVC increase, AU/year</td>
<td>188±176</td>
<td>45±25</td>
<td>138±35</td>
<td>379±175</td>
<td>&lt;0.0001</td>
</tr>
</tbody>
</table>

Results are mean±SD or number of patients (percentage). AU, arbitrary unit; AVC, aortic valve calcification; MPG, mean pressure gradient.

Clinical implications
Calcific aortic valve disease was long considered as a degenerative and, therefore, permanent condition. Clinical and histological similarities with atherosclerosis have led to the hypothesis that statins may prevent AS progression. The Scottish Aortic Stenosis and Lipid Lowering Trial, Impact on Regression (SALTIRE), SEAS and Aortic Stenosis Progression Observation: Measuring Effects of Rosuvastatin (ASTRONOMER) trials assessed the effect of statins on progression of AS in asymptomatic patients and failed to demonstrate any effect.17–19 It has thus been suggested that any effect of statins may only be observable in the early phase in patients with mild AS or even in patients with aortic valve sclerosis.22 Such a strategy seems unrealistic and the cost-effectiveness is questionable as it would require treating one-quarter of the population aged 65 years or older.23 The negative results of these trials more reasonably underline the necessity for exploring other pathophysiological pathways, such as calcium–phosphorus metabolism, vitamin D linked to bone remodelling23 or impairment of NO signalling.24 As we observed, a faster AVC progression with baseline AVC load or haemodynamic severity, if a medical strategy aimed at preventing AVC progression emerges, our study supports the need to treat all subsets of patients with AS, even those with severe AS. Additionally, our results support current guidelines recommending a closer follow-up of patients with AS, as haemodynamic severity increases as these patients are more prone to rapid progression.12

Study limitations
The present study deserves several comments. First, it was a single-centre study. Nevertheless, it was a prospective study with a wide range of AS severity and predefined intervals between visits with both simultaneous (same day) and blinded echocardiographic and CT assessment. Second, we decided to use trans-aortic MPG for the determination of AS severity as it is more robust and reproducible than AHA. However, our conclusions...
remained unchanged when AVAi or PV were used instead of MPG. Finally, the number of patients with severe AS (n=9) and patients with bicuspid aortic valve (n=24) considered in the progression analysis was relatively low. The low number of patients with severe AS is at least partially explained by the fact that only patients with at least 2 years of follow-up were enrolled in the present study. Our conclusions both in the subset of bicuspid patients and patients with severe AS deserve confirmation in larger sample sizes. Additionally, diagnosis of bicuspid aortic valve was based on echocardiography which may be difficult in case of severe calcification. Finally, we decided to enrol only participants with at least 2 years of follow-up to ensure enough time for haemodynamic or anatomic changes to occur.

CONCLUSION

In a prospective cohort of patients with a wide range of AS severity, we showed that AS progressed faster with increasing haemodynamic or anatomic severity, but there was no difference in progression related to aortic valve anatomy or gender. Our results suggest that a medical strategy aimed at preventing AV progression may be useful in all subsets of patients with AS including those with severe AS and support the recommended closer follow-up of patients with AS, as the severity of AS increases.

REFERENCES