Aortic root complications in Marfan’s syndrome: identification of a lower risk group

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Abstract

Objectives—The purpose of this study was to examine clinical and echocardiographic predictors of outcome in a cohort of patients with Marfan’s syndrome.

Background—Serial echocardiographic measurements of aortic root dimensions are an important clinical method for monitoring patients with Marfan’s syndrome. However, there are few data on the prognostic importance of echocardiographic variables for risk stratification and timing of aortic root replacement.

Methods—In 89 consecutive patients with Marfan’s syndrome (age range 1–54 years) clinical and serial echocardiographic data (n = 62) were evaluated as potential predictors of outcome (mean range follow up 4 (<1–16) years). Aortic sinus diameter and an aortic ratio normalised for age and body surface area were examined using Kaplan-Meier life table and Cox regression analysis, with the end point defined as death or surgery for ascending aortic dissection and events defined as an end point or surgery for ascending aortic aneurysm.

Results—Overall actuarial survival at two and five years was 96% and 92% and event free survival was 85% and 76%, respectively. There were five deaths due to aortic dissection, four patients survived surgery for ascending dissection, and nine underwent root replacement with a composite graft for ascending aneurysm. Those with aortic events were older (35 (12) v 25 (13) years, P = 0.007) and had greater initial aortic root dimensions (47 (14) v 33 (8) mm, P < 0.0001) and ratios (1.6 (0.5) v 1.3 (0.2), P < 0.0001). In the 62 patients with serial echocardiographic follow up, the rate of aortic root dilatation was more rapid in those with events (15 (17) v 0 (3)%/year, P < 0.0001). Utilising a Cox proportional hazards model, the groups with an initial aortic ratio ≥ 1.3 or an annual change in aortic ratio ≥ 5% had a relative risk of an aortic complication of 2.7 and 4.1, respectively (95% confidence limits 1.5 to 4.8 and 1.8 to 9.3). Only one of 31 patients with an initial aortic ratio of < 1.3 and a rate of change of < 5% had an event (five year event free survival 97%).

Conclusions—A low risk subgroup of patients with Marfan’s syndrome can be identified as those with an aortic ratio < 1.3 and an annual change in aortic ratio of < 5%. These findings are helpful in optimising echocardiographic monitoring and risk stratification of patients with Marfan’s syndrome.

Keywords: Marfan’s syndrome; aortic root complications; echocardiography; prognosis

The Marfan syndrome is an autosomal dominant inherited disorder of connective tissue with pleiotropic and variable manifestations in multiple organ systems. Aortic root pathology, including aneurysmal dilatation and dissection, is the major cause of morbidity and mortality in these patients. Serial echocardiographic aortic root measurements are an important means of surveillance, given the improved long-term prognosis after aortic root replacement.1 It was suggested in a recent study that generalized aortic root dilatation was a predictor of subsequent aortic complications in patients with Marfan’s syndrome.4 However, that study1 and two other recent studies5,6 showed wide variability in the rates of change in aortic dimensions in cohorts of patients with Marfan’s syndrome. For optimal monitoring and risk stratification, more data are needed to determine which clinical and echocardiographic variables predict outcome in patients with Marfan’s syndrome.

The aim of this study was to identify clinical and echocardiographic predictors of outcome in a large cohort of patients with Marfan’s syndrome. We hypothesised that aortic complications were more likely in those with (a) greater initial aortic root dimensions for age and body size and (b) more rapid progression of aortic dilatation.

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Patients and methods

Patients

The records of 100 patients with definite Marfan’s syndrome referred to the University of Washington Medical Center or Children’s Hospital and Medical Center genetics clinics between 1977 and 1994 were reviewed for possible inclusion in this study. Follow up information was available or could be obtained in 89 patients (from 68 families) who comprise the study group. Most individuals with Marfan’s syndrome in our geographical area are referred to these clinics, even when the primary physician is not university affiliated. In addition, the local Marfan society was contacted to ensure that all known individuals
with Marfan’s syndrome were included. The diagnosis of the Marfan’s syndrome was made according to the criteria of Pyeritz and McKusick and only patients satisfying strict diagnostic criteria were included. The mean (range) period of follow up from the initial diagnosis of Marfan’s syndrome was 4 (<1–16) years. The protocol for this study was approved by the University of Washington and Children’s Hospital Institutional Review Boards and informed consent was obtained.

FOLLOW UP
Clinical, echocardiographic, and outcome data available were collected by reviewing genetics clinic and hospital charts and telephone interview. A standardized data collection form was utilised, which included demographic details, vital statistics, whether aortic root surgery had been performed and a detailed account of the patient’s usage of β blocking medication. The defined outcomes were (a) the end point of death or root replacement for ascending dissection or (b) ascending aortic events defined as those with an end point plus those undergoing surgery for ascending aortic aneurysm. The decision to perform aortic surgery was made by each patient’s physician based on clinical, as well as echocardiographic, data. The aortic root ratio or the change in root ratio was not used in clinical decision making. In addition to aortic root dilatation, the indications for aneurysm surgery included increasing aortic regurgitation or new cardiac symptoms, or both, in seven (78%) of nine of these patients.

There were four patients with descending aortic dissections of whom only one was considered to have reached an end point (one also had aortic root replacement for ascending aneurysm). The other three patients with isolated descending aortic dissections were censored at the time of the event but were not considered an end point, as the study goal was to predict ascending aortic complications. One patient died after descending aortic dissection. Inclusion of this death as an end point does not change the statistical significance of the results. Patients were also grouped into those receiving β blocking drugs for >12 months and those who never received them or took them for <12 months.

ECHOCARDIOGRAPHIC FOLLOW UP
Echocardiographic data were available in 80 (90%) of 89 patients. Serial echocardiographic studies were available in 62 (70%) of 89 patients with a median (range) of serial studies per patient of 4-6 (2-13). Patients who had not had an echocardiogram within the previous year underwent a clinical evaluation and echocardiographic examination focusing on aortic dimensions and pathology at the University of Washington Medical Center (n = 18).

Detailed measurements of the aortic root were made in the parasternal long axis view according to the method of Roman et al by an investigator unaware of clinical outcome in the 98 echocardiograms in which the original video tape could be obtained. In the other studies (n = 199) official reports were used. Aortic diameter measurements were made using the leading edge technique at end diastole (defined as the peak of the R wave on the electrocardiogram). Maximal diameter measurements were recorded at the aortic annulus, the sinuses of Valsalva, the sinotubular junction, and proximal ascending aorta 1–2 cm above the sinotubular junction. All diameters were measured perpendicular to the long axis of the aorta.

The maximum dimension of the sinuses of Valsalva (available in all studies) was used to calculate an aortic ratio. The observed dimension was divided by the predicted dimension based on age and body surface area using the following established regression formulas derived from normal individuals:

Children (age < 18 years): predicted sinus dimension (cm) = 1.02 + (0.98 × BSA (m²))
Adults (age 18–40 years): predicted sinus dimension (cm) = 0.97 + (1.12 × BSA (m²))
Adults (>40 years): predicted sinus dimension (cm) = 1.92 + (0.74 × BSA (m²))

where BSA is body surface area.

This allowed for comparisons of aortic root dimensions among all patients regardless of age and body size. For example, an aortic sinus ratio of 1.3 indicates a 30% enlargement of the aortic root above the mean for that patient’s age and body surface area. The rate of aortic root dilatation normalized for age and body size was calculated by determining the change in aortic ratio between the final and initial echocardiograms and dividing by the interval between them in years. Mitral valve prolapse and aortic regurgitation were evaluated using established cross sectional echocardiographic and colour Doppler diagnostic criteria.8-10

STATISTICAL ANALYSIS
Demographic and echocardiographic variables in patients with versus those without an aortic event or end point were compared using χ² analysis for dichotomous variables and the unpaired Student’s t test for continuous variables. Data are presented as mean (one SD). Kaplan-Meier life table analysis was performed for overall and event free survival. The log rank test was used to examine differences between survival curves in those who had taken β blocking medication for > 1 year or ≤ 1 year. The time period of one year for β blocker treatment was chosen because some patients took β blockers for only a few days and then discontinued them because of side effects. The results are not affected by using a different cut off point (e.g. one month) for duration of β blocker treatment.

Cox proportional hazards regression was performed to examine baseline variables that were found on univariate analysis to be associated with a poor outcome. For the Cox regression model, an aortic ratio > or < 1.3, and an
Table 1: Demographic and echocardiographic features of 89 patients with Marfan syndrome

<table>
<thead>
<tr>
<th>Feature</th>
<th>Value (SD) (range)</th>
<th>n (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean (SD) (range) age at entry (years)</td>
<td>21 (13) (1-54)</td>
<td>89</td>
</tr>
<tr>
<td>Age &lt; 18 years at enrolment (%)</td>
<td>14</td>
<td>64</td>
</tr>
<tr>
<td>Sex (M/F)</td>
<td>49/40</td>
<td></td>
</tr>
<tr>
<td>Family history (%)</td>
<td>56</td>
<td></td>
</tr>
<tr>
<td>Organ system (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cardiovascular</td>
<td>91</td>
<td></td>
</tr>
<tr>
<td>Musculoskeletal</td>
<td>94</td>
<td></td>
</tr>
<tr>
<td>Ocular</td>
<td>83</td>
<td></td>
</tr>
<tr>
<td>Clinical findings (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mitral prolapse</td>
<td>63</td>
<td></td>
</tr>
<tr>
<td>Spinal deformity</td>
<td>38</td>
<td></td>
</tr>
<tr>
<td>Hyperextensibility</td>
<td>54</td>
<td></td>
</tr>
<tr>
<td>Ectopia lentis</td>
<td>44</td>
<td></td>
</tr>
<tr>
<td>&amp;blocker treatment (%)</td>
<td>≥ 1 year 36 &lt; 1 year 64</td>
<td></td>
</tr>
<tr>
<td>Aortic regurgitation (%)</td>
<td>None 71 Mild to moderate 11 Severe 3 Unknown 15</td>
<td></td>
</tr>
<tr>
<td>Mitral regurgitation (%)</td>
<td>None 59 Mild to moderate 25 Severe 1 Unknown 15</td>
<td></td>
</tr>
</tbody>
</table>

Results

DEMOGRAPHIC FEATURES

Table 1 lists the demographic and clinical characteristics of the 89 patients with Marfan’s syndrome. There were 49 males and 40 females with an age range from 1 to 54 (median 21) years. Cardiovascular and musculoskeletal features were present in nearly all patients. Ocular findings were present in 83% of patients and included iridodonesis, lens dislocation, or a history of retinal detachment. A positive family history of Marfan’s syndrome was present in 56% of patients, while 44% represented new mutations. Although 45% (37 of 83 patients) of patients had taken &beta; blocking medication at some time in the follow-up period, only 36% (30 of 83) took &beta; blockers continuously for greater than 12 months. &beta; blocker history was unknown in six patients.

OVERALL SURVIVAL AND EVENT-FREE SURVIVAL

Figure 1 shows Kaplan-Meier life table analysis for overall survival, freedom from aortic dissection, and event free actuarial survival. Actuarial survival at two and five years for the whole group was 96% and 92%, and freedom from all events was 85% and 76%, respectively. The Kaplan-Meier mean total survival was 14-6 (95% confidence intervals CI 13-4 to 15-8) years and mean event free survival was 12-0 (95% CI 10-3 to 13-6) years from presentation.

At follow up 71 of 89 patients were alive with no ascending aortic complications. The 18 end points included five deaths due to ascending dissection, four patients who survived emergency aortic root replacement for ascending aortic dissection, and nine patients who underwent a composite graft replacement for ascending aortic aneurysm (table 2). Surgery or death occurred < one year after the initial diagnosis in 11 of 18 patients with an end point, thus, follow up is shorter in those with an event. The mean (range) age at death was 30 (17-54) years and for all aortic complications was also 30 (10-54) years.

CLINICAL CHARACTERISTICS OF THOSE WITH AND THOSE WITHOUT AORTIC COMPLICATIONS

There were no significant differences between those with and those without aortic complications other than an older age and shorter follow up period in those with aortic complications (table 3). Fifty three per cent of those having an aortic complication had a first degree relative with an aortic complication or died due to aortic dissection. Results are similar if only those with a definite end point (death or dissection) are compared with the remainder of the study group.

ECHOCARDIOGRAPHIC FOLLOW UP

In the 80 patients with one or more echocardiograms, the initial aortic sinuses dimension in those with an aortic complication averaged 47 (range 31-83) mm compared with 33 (range 20-50) mm in those without a complication (P < 0.0001). This was reflected in the aortic ratio which averaged 1:6 in those with an event compared with 1:3 in those without (P < 0.0001). The initial aortic ratio was ≥ 1:3 in 77% of patients with a complication versus 35% of patients without (χ² = 0.01). Similar changes were seen in the aortic dimensions at
the sinotubular junction and ascending aortic levels, although the number of measurements was smaller. The mean sinotubular junction dimension on the final echocardiogram in those with versus those without an aortic complication was 41 (10) v 31 (5) mm (P = 0.001). Similarly, the ascending aorta measured 35 (11) v 27 (7) mm (P = 0.03). Although the average sinus of Valsalva dimension on the most recent echocardiographic study before the aortic complication was 51 (range 38–83) mm, it is noteworthy that four patients (all women) had an aortic root dimension of only between 38 and 46 mm in the year before aortic dissection. Only one of these women was postpartum (fig 2).

Table 4 compares echocardiographic findings in the 62 patients with serial studies. The mean baseline sinus dimension and aortic ratio were no different in those who developed a complication versus those who remained event free. However, the changes in aortic sinus dimension and aortic ratio/year of follow up were significantly greater in those with a complication than in those without. The annual rate of change in aortic ratio was ≥5% in only four (8%) of 51 of those without a complication compared with five (83%) of six with a complication (χ² = 0.001) (table 4). A more rapid rate of dilatation (i.e. ≥5%/year) was detected only in patients who were subsequently referred for aortic root replacement for ascending aneurysm. Serial echocardiographic data were available only in one patient with acute ascending dissection.
Table 4  Serial echocardiographic findings (n = 62)

<table>
<thead>
<tr>
<th>Event (n = 6)</th>
<th>No event (n = 56)</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Body surface area</td>
<td>1.6 (0.5)</td>
<td>1.9 (0.3)</td>
</tr>
<tr>
<td>Sinus dimension (mm) (n = 62)</td>
<td>34 (8)</td>
<td>37 (6)</td>
</tr>
<tr>
<td>Baseline</td>
<td>38 (7)</td>
<td>57 (10)</td>
</tr>
<tr>
<td>Follow up</td>
<td>5 (5)</td>
<td>20 (9)</td>
</tr>
<tr>
<td>A ratio</td>
<td>0.7 (1)</td>
<td>5 ± 6</td>
</tr>
<tr>
<td>A ratio/year</td>
<td>1.3 (0.2)</td>
<td>1.2 (0.2)</td>
</tr>
<tr>
<td>Follow up</td>
<td>1.3 (0.2)</td>
<td>1.8 (0.4)</td>
</tr>
<tr>
<td>A ratio</td>
<td>0.03 (0.1)</td>
<td>0.6 (0.3)</td>
</tr>
<tr>
<td>A ratio/year</td>
<td>0.00 (0.03)</td>
<td>0.15 (0.17)</td>
</tr>
</tbody>
</table>

Values are mean (SD). NS, not significant.

Figure 3 Examples of the rate of change in aortic sinus ratio over the follow up period. In these 12 patients with Marfan's syndrome, two underwent surgery for ascending aortic aneurysm and in increasing aortic regurgitation (A) while the remaining 10 had no aortic complications (B).

Figure 4 Initial aortic sinus ratio versus age showing that there is no significant relation between these two variables. Points with no event (○), those who died (■), and those who had ascending aortic surgery for dissection (△) or aneurysm (□) are indicated.

A lower risk group was defined as patients with an initial aortic ratio < 1.3 and a rate of change of aortic ratio < 5%/year. Only one (3%) of 31 of these patients had an aortic complication compared with 14 (36%) of 39 patients who had an initial aortic ratio of ≥ 1.3 or an annual rate of change of ≥ 5%. The relative risk of a complication in this low risk subgroup was 0.3 (95% CI 0.09 to 0.71, \( \chi^2 = 0.001 \)). The predictive value of these variables in predicting freedom from an event was therefore 97%. Dissection or death occurred in only one (3%) of 31 of this low risk subgroup versus six (15%) of 39 with an aortic ratio ≥ 1.3 or annual rate of change ≥ 5%. Use of this more definitive end point did not reach statistical significance because of the small number of end points.

To examine the effect of patients who presented acutely on the Cox regression model, the eight patients with follow up times < one month were excluded from analysis. Similar results were obtained with a relative risk of a complication of 2.8 (95% CI 1.4 to 5.5) for those with an initial aortic ratio of ≥ 1.3 and a relative risk of 5.9 (95% CI 2.0 to 17.3) for those with a rate of change in aortic ratio of ≥ 0.05/year.

β BLOCKER TREATMENT

Thirty patients received β blockers for ≥ one year while 53 patients either never received β blockers or took them for < one year. Aortic ratios and the change in ratio between initial and final echocardiograms did not differ between these groups. Actuarial freedom from all events was no different in those receiving β blockers (69%, nine events in 28 patients) compared with those not receiving β blockers (85%, eight events in 55 patients) at five years (P = not significant, log rank test).
Discussion
ECHOCARDIOGRAPHIC PREDICTORS OF OUTCOME
This study demonstrates that echocardiographic variables are powerful predictors of outcome in patients with Marfan's syndrome. In those in whom serial echocardiographic measurements of aortic dimensions were available, the rate of dilatation of the aortic root was significantly more rapid in those with complications. The mean change in aortic ratio in those suffering a complication was 0.15/year indicating an average increase of 15% in aortic dimension/year, after allowing for the effect of increasing age and body size. This contrasts with almost no change in the aortic ratio seen in those not experiencing a complication. Thus, an annual change in the aortic ratio < 5% indicates that the patient is unlikely to suffer an aortic complication (predictive value 92%). In an individual patient with Marfan's syndrome, a consistent pattern of change on serial echocardiography is helpful in distinguishing progressive root dilatation from measurement variability.

Although the number of serial echocardiographic studies in patients with a complication was small because a number of patients had their complication soon after their initial study, our findings remain significant even when these patients are excluded. Thus, the finding of more rapid dilatation underscores the need for more frequent surveillance especially in patients at high risk, either because of a strong family history of aortic complication, or significant aortic dilatation at baseline.

The importance of calculating an aortic ratio also must be emphasized. Aortic dimensions were between 38 and 46 mm in four patients recorded within one year of ascending dissection. These four patients (all women) had body surface areas between 1.70 and 2.27 m², so that their aortic ratios were between 1.3 and 1.4. As women generally have a smaller body size, the magnitude of aortic dilatation may not be appreciated when only the absolute sinus dimension is considered. The prognostic value of the aortic ratio is further emphasised by the Cox proportional hazards model. Patients with an initial aortic sinus ratio > 1.3 had an almost threefold risk of an aortic complication over the follow up period compared with that in those with an aortic ratio < 1.3. In a man with a body surface area of 2.0 m², an aortic ratio of 1.3 corresponds to an aortic sinus dimension of approximately 42 mm. In a woman with a body surface area of 1.8 m², a ratio of 1.3 corresponds to an aortic sinus dimension of 39 mm. The current general recommendation for prophylactic aortic root replacement is at an aortic dimension of 55–60 mm,\(^\text{11,15}\) which would correspond to an aortic ratio of 1.7–1.9 in the male and 1.8–2.0 in the female examples given here.

In our population a substantial number of complications occurred in patients with aortic root dimensions less than that for which prophylactic root replacement is routinely performed (fig 2) as noted in other studies.\(^\text{11,15}\) However, no complications occurred in patients with an aortic ratio < 1.3.

LONG-TERM OUTCOME IN PATIENTS WITH MARFAN'S SYNDROME
With current cardiac surgical techniques and improved echocardiographic monitoring, long-term survival in patients with Marfan's syndrome is reasonably good. However, there is a substantial incidence of aortic complications. All deaths in this series were due to aortic dissection, including two patients not previously known to have Marfan syndrome or aortic root pathology. While the 20% incidence of aortic complications over a mean follow up period of four years is similar to that in recently published series,\(^\text{1,3}\) the survival of 92% for the whole cohort at five years seems to be better, probably because all patients underwent a composite graft usually with the classic Bentall's technique\(^\text{15}\) rather than the older technique of an interposition tube graft. There were no early postoperative deaths with elective surgery (n = 9) for aortic dilatation. With emergency surgery for dissection, four (50%) of eight patients died intraoperatively or in the early postoperative period. The long-term survival of patients with aortic complications who survived surgery was excellent, with no known deaths a mean of 8.1 years after aortic root replacement. However, four of 13 patients surviving aortic surgery required reoperation, two for further dissection, one because of graft infection, and one because of graft failure. These data are in agreement with previous studies showing a five and 10 year actuarial survival after composite graft repair between 71 and 93% and 54 and 76%, respectively.\(^\text{11,15}\)

OTHER PREDICTORS OF OUTCOME
The importance of factors not evaluated in this study cannot be discounted. For example, the high incidence of complications in patients who had first degree relatives with aortic complications emphasises the adverse prognostic importance of a positive family history of aortic dissection.\(^\text{16,19}\) In addition, cardiovascular risk may differ within families as well as with some members having milder phenotypes not associated with aortic involvement.\(^\text{20}\)

The retrospective nature and relatively small sample size of this study make interpretation of the data pertaining to \(\beta\) blocker treatment difficult. Some patients presented acutely with dissection and previously undiagnosed Marfan's syndrome, a bias that favours \(\beta\) blockade. Conversely, it is likely that those with more severe disease (e.g. family history, larger aortic dimension, and rapid rate of aortic dilatation) were selected for treatment which would bias against \(\beta\) blockade. Overall, no significant differences between the two groups were observed.

Conclusions
Long-term prognosis in patients with Marfan's syndrome is reasonably good, but a significant number of aortic complications occur, even in patients with only moderately dilated aortic roots. Until a molecular based approach is widely available that will identify patients at high cardiovascular risk, echocardiographic
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variables will remain the most important prognostic indicators. The present study suggests that any patient with an aortic ratio $\geq 1.3$ (30% greater than normal) or a change in aortic ratio $>5\%$/year should be followed closely as there is a high risk of complications. Patients with aortic ratios of $<1.3$ and a rate of change of $<5\%$ seem to be at low risk. Prospective validation of these proposed criteria is needed.

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