Patients with severe pulmonic stenosis (PS) have right ventricular (RV) diastolic filling abnormalities detectable by tricuspid valve pulsed Doppler examination. To determine if these abnormalities persist long term after successful therapy of PS, 19 patients were examined 8 ± 3 years after PS therapy. At the time of follow-up Doppler examination, the PS gradient was 15 ± 8 mm Hg. From the tricuspid valve inflow Doppler study, the following measurements were obtained at peak inspiration: peak velocities at rapid filling (peak E) and during atrial contraction (peak A), ratio of peak E to peak A velocities, RV peak filling rate normalized for stroke volume, deceleration time, the fraction of filling in the first 0.33 of diastole as well as under the E and A waves, and the ratio of E to A area. Data from PS follow-up patients were compared with our previously reported data from 12 age-related control subjects and 14 untreated patients with PS. Patients with PS who were followed up had higher peak E velocity (0.75 ± 0.14 vs 0.59 ± 0.21 m/s), lower peak A velocity (0.47 ± 0.09 vs 0.64 ± 0.28 m/s), higher E/A velocity ratio (1.65 ± 0.33 vs 1.11 ± 0.52), higher 0.33 area fraction (0.52 ± 0.08 vs 0.34 ± 0.14), lower A area fraction (0.29 ± 0.06 vs 0.45 ± 0.21) and higher E/A area ratio (2.48 ± 0.82 vs 1.73 ± 1.06) than PS patients without treatment (p < 0.03). All Doppler indexes of the patients with PS who were followed up were the same as those of the control subjects except for the peak E velocity that was slightly higher (0.75 ± 0.14 vs 0.63 ± 0.11 m/s), the peak A velocity that was slightly higher (0.47 ± 0.09 vs 0.38 ± 0.09 m/s) and the E/A area ratio that was slightly lower (2.48 ± 0.82 vs 3.50 ± 1.25) (p < 0.03). Thus, at long-term follow-up, all RV diastolic filling indexes in successfully treated patients with PS improved compared with the untreated patients and approached values found in normal subjects. These data suggest that RV diastolic filling abnormalities in patients with PS are reversible over the long term and are therefore probably related to hypertrophy rather than fibrosis and scarring. (Am J Cardiol 1991;68:648–652)

In recent years, pulsed Doppler echocardiography has been used to assess right ventricular (RV) diastolic filling in a variety of diseases including valvular pulmonic stenosis (PS), pulmonary hypertension, constrictive pericarditis and cardiac tamponade.1-4 From the tricuspid valve Doppler recording, peak flow velocities, filling rates, and the proportion of filling in the various phases of diastole can be measured for the right ventricle.5 Normal values for the tricuspid valve Doppler indexes have been reported for the fetus, newborn infant and child.6-10 Recently, we reported the use of these Doppler indexes to detect abnormal patterns of RV diastolic filling in children with PS.4 Children with PS had a decreased percentage of the total Doppler area in the first third of diastole and an increased percentage of the total Doppler area under the A wave suggesting a relative shift of RV filling to late diastole. Furthermore, these diastolic filling abnormalities did not improve immediately after successful relief of the RV outflow obstruction, suggesting that afterload mismatch was not the direct cause of the observed diastolic filling abnormalities.

In this study, we hypothesized that RV hypertrophy is the cause of the impaired RV early diastolic relaxation found in children with valvular PS. Furthermore, if the RV diastolic filling abnormalities were caused by hypertrophy alone rather than fibrosis or scarring, then these abnormalities should return to normal as hypertrophy regresses over the long-term follow-up period after successful relief of PS. To test this hypothesis, we
assessed RV diastolic filling using pulsed Doppler echocardiography in 19 patients who were examined 8 ± 3 years after successful relief of PS.

METHODS

Patients: The study included 19 patients who were examined 8 ± 3 years (mean ± standard deviation) after successful relief of PS (Table I). These patients were randomly selected from all patients with PS undergoing routine follow-up evaluation in the outpatient clinic. Selection criteria included: (1) evidence by Doppler examination of successful relief of PS (peak gradient <25 mm Hg), (2) absence of additional congenital defects such as tricuspid stenosis or left-to-right shunts that might alter the tricuspid valve Doppler recording, and (3) absence of significant tricuspid regurgitation that might mask RV diastolic filling abnormalities.

The PS follow-up group was 5 to 19 years old (mean 10.9) and weighed 18 to 83 kg (mean 45.2). Ten patients had previous pulmonary balloon valvuloplasty and 9 had surgical pulmonary valvotomy. Ten patients were treated before and 9 were treated after the age of 2 years. Data from PS patients at long-term follow-up were compared with our previously reported data from 12 age-related control subjects and 14 untreated patients with severe PS (Table I).

Echocardiographic examinations: All study participants underwent a complete 2-dimensional and Doppler echocardiographic examination with the use of a 128-element phased-array ultrasound system and a variety of transducers appropriate for patient size. Tricuspid valve Doppler examinations were obtained from the parasternal short-axis or apical 4-chamber view. The sample volume was positioned so as to record the maximal velocities through the valve (usually near the tips of the leaflets). Based on prior studies with use of simultaneous thermister and tricuspid valve Doppler recordings, the velocities through the tricuspid valve vary significantly throughout the respiratory cycle with maximal velocities occurring at peak inspiration. Therefore, to obtain all Doppler measurements at a standard time in the respiratory cycle, only beats recorded at peak inspiration were used.

From the Doppler spectral recordings, the peak velocities during rapid ventricular filling (peak E) and during atrial contraction (peak A) were measured, and the ratio of peak E to peak A velocities was calculated. To determine the Doppler pattern of RV filling, several areas under the Doppler tracing were measured using previously described methods: (1) the total area under the velocity envelope throughout diastole; (2) the area under the velocity curve for the first 33% of diastole (0.33 area); (3) the E area, or the triangular area formed by extrapolating a straight line down from the peak E velocity to the baseline; and (4) the A area, or the triangular area formed in a similar manner under the peak A velocity. To determine the percentage of the total velocity envelope occupied by the individual areas, the area of filling fractions were calculated as the individual areas divided by the total area under the Doppler tracing. Also, the ratio of E and A areas was calculated. The peak filling rate normalized for stroke volume was calculated as the peak E velocity divided by the total velocity time integral, and the deceleration time was measured from the peak E velocity to the time when the Doppler curve returned from the peak E velocity to the baseline.

Color, pulsed and continuous-wave Doppler examinations of the pulmonary valve were obtained from parasternal or subcostal views. With use of the view that provided the highest value for the peak velocity of the PS jet, the peak instantaneous pressure gradient across the pulmonary valve was calculated from the Bernoulli equation. Pulmonary insufficiency was diagnosed using color flow imaging techniques and was considered to be significant if the M-mode echocardiographic examination showed evidence of paradoxical septal motion or RV end-diastolic dimension >95% for body surface area, or both.

All Doppler examinations were recorded at a paper speed of 100 mm/s. The Doppler areas were traced from the paper recording using a digitizing tablet with a crosswire cursor, a personal computer and commercially available computer software. The Doppler velocities and areas were measured by tracing the outermost border of the spectral recordings.

Statistical analysis: Three cardiac cycles were measured and averaged to obtain each Doppler value. Statistical comparisons between the PS follow-up patients and the control subjects and between the untreated PS patients and the PS follow-up patients were obtained using unpaired t tests and Bonferroni's correction for multiple comparisons. To determine if differences in RV diastolic filling patterns existed between different patients in the PS follow-up group, the group was divided into several subgroups which were then compared using unpaired t tests. Subgroup analyses included: (1) comparison of PS follow-up patients >2 years old at

| TABLE I Demographic Data for the Three Patient Groups |
|-----------------|-----------|-----------|-----------|
|                  | Control   | PS F/U    | PS Pre    |
| No. of patients  | 12        | 19        | 14        |
| Age (years)      | 8.6 (4.5–16) | 10.9 (5–19)* | 5.1 (0.4–18)† |
| Weight (kg)      | 32.7 ± 15.4 | 45.2 ± 21.7* | 21.6 ± 20† |
| Height (cm)      | 133 ± 20   | 145 ± 20*  | 103 ± 32†  |

* p <0.01 compared with PS pre; †p <0.01 compared with control subjects. F/U = follow-up; Pre = untreated; PS = pulmonary stenosis.
As a result, the E/A velocity ratio of PS follow-up patients was higher than that of the untreated PS patients (p = 0.05) and the same as that of the control subjects. The A area fraction of the PS follow-up patients had decreased significantly compared with that of untreated PS patients and was the same as that of normal subjects. As a result, the E/A area ratio of the PS follow-up group was significantly higher than that of untreated PS patients but did not quite reach the value found in normal subjects.

The deceleration time of the PS follow-up patients was longer than that found in the untreated PS patients but the same as that found in normal subjects. The normalized peak filling rates of the 3 patient groups were not significantly different.

SUBGROUP ANALYSIS: To determine the effect of age at the time of treatment on the observed Doppler findings, the PS follow-up group was divided into 2 subgroups: (1) patients >2 years old at the time of therapy (n = 10), and (2) patients <2 years old at the time of therapy (n = 9). No differences were found in any diastolic filling indexes between the 2 subgroups.

To determine the effect of the type of therapy on the observed Doppler findings, the PS follow-up group was divided into 2 subgroups: (1) patients treated with surgery (n = 9), and (2) patients treated with balloon valvuloplasty (n = 10). No differences were found in any of the RV diastolic filling indexes between the 2 subgroups.

To determine if significant pulmonary insufficiency had an effect on the observed Doppler findings, a subgroup of PS follow-up patients was formed by excluding all patients with significant pulmonary insufficiency and RV diastolic dimension >95% for body surface area. A two-tailed p value <0.03 was used to indicate a significant intergroup difference. All values are mean ± standard deviation.

**RESULTS**

Patients: The PS follow-up patients did not differ from the control subjects in age, height, weight or heart rate. Compared with untreated PS patients, the PS follow-up patients were older, had greater heights and weights, and had slower heart rates. Before treatment, the PS follow-up patients had the same pulmonary valve gradient as the untreated PS patients (67 ± 18 vs 71 ± 35 mm Hg, p = 0.66). After treatment, the PS follow-up patients had a significantly lower pulmonary valve gradient (15 ± 8 mm Hg).

**Echoangiographic studies: Comparisons of Pulmonic Stenosis Follow-up Patients with Control Subjects and Untreated Pulmonic Stenosis Patients:** Mean values for the Doppler measurements of the 3 patient groups are listed in Table II. The peak E velocity of the PS follow-up patients was significantly higher than that of the untreated PS patients and the control subjects. The peak A velocity of the PS follow-up patients was significantly lower than that of the untreated PS patients but still higher than that of control subjects. As a result, the E/A velocity ratio of PS follow-up patients was increased to a value not different from that found in normal subjects.

Compared with untreated PS patients, the 0.33 area fraction of the PS follow-up patients had increased significantly to a value not different from that found in control subjects. Likewise, the E area fraction of the PS follow-up patients was higher than that of the untreated PS patients (p = 0.05) and the same as that of the control subjects. The A area fraction of the PS follow-up patients had decreased significantly compared with that of untreated PS patients and was the same as that of normal subjects. As a result, the E/A area ratio of the PS follow-up group was significantly higher than that of untreated PS patients but did not quite reach the value found in normal subjects.

The deceleration time of the PS follow-up patients was longer than that found in the untreated PS patients but the same as that found in normal subjects. The normalized peak filling rates of the 3 patient groups were not significantly different.

**DISCUSSION**

In a previous study, we showed that children with severe PS have RV diastolic filling abnormalities detectable with Doppler echocardiography. In these children, the tricuspid Doppler examination showed a decreased percentage of the total Doppler area in the first third of diastole and an increased percentage of the total Doppler area under the A wave, suggesting a relative shift of RV filling to late diastole. This abnormal tricuspid Doppler pattern resembles the mitral Doppler pattern observed by Appleton et al in patients with impaired left ventricular early diastolic relaxation and normal left ventricular filling pressures. Immediately after balloon valvuloplasty and successful relief of the high afterload, RV diastolic filling abnormalities per-

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### TABLE II Tricuspid Valve Doppler Measurements

<table>
<thead>
<tr>
<th>Measurement</th>
<th>Control Subjects</th>
<th>PS F/U</th>
<th>PS Pre</th>
</tr>
</thead>
<tbody>
<tr>
<td>Peak E (m/s)</td>
<td>0.63 ± 0.11</td>
<td>0.75 ± 0.14*</td>
<td>0.59 ± 0.21</td>
</tr>
<tr>
<td>Peak A (m/s)</td>
<td>0.38 ± 0.09</td>
<td>0.47 ± 0.09**</td>
<td>0.64 ± 0.28*</td>
</tr>
<tr>
<td>E/A vel.</td>
<td>1.74 ± 0.51</td>
<td>1.65 ± 0.33†</td>
<td>1.11 ± 0.52‡</td>
</tr>
<tr>
<td>Total VTI (ms)</td>
<td>0.12 ± 0.02</td>
<td>0.13 ± 0.02</td>
<td>0.12 ± 0.03</td>
</tr>
<tr>
<td>0.33 area fx</td>
<td>0.51 ± 0.12</td>
<td>0.52 ± 0.08†</td>
<td>0.34 ± 0.14*</td>
</tr>
<tr>
<td>E area fx</td>
<td>0.71 ± 0.08</td>
<td>0.67 ± 0.07</td>
<td>0.57 ± 0.19*</td>
</tr>
<tr>
<td>A area fx</td>
<td>0.24 ± 0.10</td>
<td>0.29 ± 0.06†</td>
<td>0.45 ± 0.21*</td>
</tr>
<tr>
<td>E/A area</td>
<td>3.50 ± 1.25</td>
<td>2.48 ± 0.82**</td>
<td>1.73 ± 1.05*</td>
</tr>
<tr>
<td>Decel. time (s)</td>
<td>0.14 ± 0.02</td>
<td>0.12 ± 0.02†</td>
<td>0.09 ± 0.04*</td>
</tr>
<tr>
<td>PFR/SV (W/s)</td>
<td>5.23 ± 0.56</td>
<td>5.79 ± 1.08</td>
<td>5.28 ± 1.96</td>
</tr>
<tr>
<td>Heart rate (beats/min)</td>
<td>77 ± 13</td>
<td>75 ± 13†</td>
<td>99 ± 27*</td>
</tr>
</tbody>
</table>

* p < 0.03 compared with control subjects; †p < 0.03 compared with PS pre.

Values are mean ± standard deviation.

A = peak velocity at atrial contraction; E = peak velocity in early diastole; Decel. = deceleration; fx = fraction; F/U = follow-up; PFR/SV = peak filling rate normalized to stroke volume; Pre = untreated; PS = pulmonary stenosis; vel. = velocity; VTI = velocity time integral.
sisted, suggesting that afterload mismatch was not the cause of the filling abnormalities. In this long-term follow-up study, all RV diastolic filling indexes in successfully treated PS patients improved compared with untreated PS patients, and approached values found in normal subjects. These data suggest that RV diastolic filling abnormalities in PS patients are reversible and are, therefore, probably related to hypertrophy rather than fibrosis and scarring. RV mass was not assessed owing to the lack of an accurate means of measuring it noninvasively; however, all untreated PS patients had echocardiographic evidence of severe RV hypertrophy while the PS follow-up patients had little or no evidence of RV hypertrophy.

Possible mechanisms of diastolic filling abnormalities Courtois et al.15 recently showed that a pattern of diastolic apex to inflow pressure gradients exists in the right ventricle during early and late diastole, similar to that reported in the left ventricle.16 In the right ventricle, however, the lowest early diastolic pressures are usually recorded in the outflow tract rather than in the apex. The form and timing of the regional ventricular pressure gradients found in their study suggest that mechanical suction of blood into the ventricular cavity is the primary mechanism of RV filling in early diastole. Mechanisms that probably contribute to mechanical suction include downward motion of the right ventricle during systole, active contraction of muscle fibers below equilibrium and resultant storage of elastic energy, and end-systolic deformation of the walls of the RV outflow tract. In the latter mechanism, blood continues to leave the ventricle after contraction has ended, thus causing the shape of the RV outflow tract to be distorted, elastic energy to be stored in the myocardium, and the walls of the outflow tract to recoil in early diastole. The importance of this mechanism is supported by the finding of the lowest early diastolic pressure in the RV outflow tract and the observation that significant narrowing of the outflow tract occurs at end-systole.15

In untreated PS patients, severe RV hypertrophy can lead to reduced end-systolic deformation of the RV outflow tract, less elastic recoil in early diastole, and a higher minimum RV pressure in early diastole. With a higher minimum RV diastolic pressure, the early diastolic pressure gradient and, thus, the peak E velocity are decreased as was observed in our untreated PS patients. At long-term follow-up, the return of the peak E velocity and the percent filling in early diastole toward normal values suggests that the end-systolic deformation of the outflow tract walls is restored as RV hypertrophy regresses.

In normal subjects, the tricuspid deceleration time is longer than the mitral deceleration time, suggesting that the thin-walled right ventricle is a less effective decelerator than the thicker-walled left ventricle.15 In untreated PS patients, we found a shortened tricuspid deceleration time compared with age-related normal subjects. It is likely that the thick-walled right ventricle of untreated PS patients quickly generates a reverse pressure gradient of sufficient magnitude to decelerate flow in early diastole. As hypertrophy regresses after successful relief of PS, the thin-walled right ventricle requires a longer period of time to generate a reverse pressure gradient, and thus the increase in deceleration time to normal values in our follow-up patients.

Factors affecting the tricuspid valve Doppler indexes: Diastolic indexes of RV relaxation can be influenced by several factors including age, heart rate, respiration and RV loading conditions. Throughout childhood, the tricuspid Doppler indexes are independent of age beyond the neonatal period5-8 and, thus, it is unlikely that age contributed to the observed improvement in RV diastolic filling. In several recent studies, tricuspid Doppler indexes have been unrelated or only weakly related to heart rate.17,18 In this study, the improved RV filling cannot be explained on the basis of heart rate since the heart rates of the control and PS follow-up patients were not different. From expiration to inspiration in normal children, the tricuspid peak E velocity increases by 26%, the peak A velocity increases by 18%, and the E/A velocity ratio remains unchanged.8 In this study, the effects of respiration were eliminated by measuring only beats at maximal inspiration. The presence of tricuspid or pulmonary insufficiency may alter the early diastolic transvalvular pressure gradient and, thus, affect the tricuspid Doppler indexes. In this study, no patient had significant tricuspid insufficiency, and exclusion of 4 patients with pulmonary insufficiency did not alter the results.

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REFERENCES


