Postoperative Outcome of Patients with Prosthetic Valve Leak

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Summary

Background: Prosthetic valve leak is a possible complication of surgical valve replacement. Although uncommon, its consequences may be serious. Few studies correlate the degree of prosthetic valve leak with clinical events.

Objective: To compare the postoperative outcome of patients with mild/moderate (Mi/Mo) or severe (Sev) prosthetic valve leak

Methods: A total of 185 patients with prosthetic valve leak were selected among 1350 patients undergoing heart valve surgery between 1999 and 2001. Of these, a sample of 58 patients (37 men) with prosthetic valve leak (36 with Mi/Mo versus 22 with Sev leak) in the pre and/or postoperative period of heart valve replacement had complete medical record data, so their clinical, laboratory and echocardiographic data could be retrospectively assessed.

Results: The incidence of reoperation was 11.1% in the Mi/Mo group, versus 22.7% in the Sev group (odds ratio = 2.35 [95% CI 0.56-9.94]). Endocarditis was the cause of reoperation in 75% of the patients of the Mi/Mo group and in 60% of the Sev group. Aortic bioprostheses were those most frequently related to leak (55.8% in the Mi/Mo group and 57.7% in the Sev group). Forty percent of the patients with previous Mi/Mo leak did not present prosthetic valve leak on postoperative day 2 versus 21.4% of the patients with Sev prosthetic valve leak. No significant differences were found regarding laboratory variables.

Conclusions: (1) Patients with severe leak are more likely to undergo reoperation. (2) Endocarditis was the most frequent cause of reoperation for any leak degree. (3) Severe prosthetic valve leak is more difficult to fully resolve after surgical treatment.

Key Words: Heart valve prosthesis; cardiac surgical procedures; endocarditis.

Introduction

Prosthetic valve leak is a relatively rare complication in patients undergoing heart valve replacement, whether for biological or mechanical prostheses. The leak may be central, paravalvular or both. Mild leak, with a low risk for pathologic consequences, may be found in a large proportion of heart valve prostheses. Provided no hemodynamic changes occur, mild leaks are not clinically significant.

Genoni et al1 found a 12.1% prevalence of leak among 598 patients, and the symptoms most commonly related to this condition were fatigue (67%), vertigo (55%), and functional class III/IV dyspnea (38%).

Jindani et al’s study2, in turn, found a 2.5% prevalence of prosthetic valve leak among a total of 1,175 prostheses (735 of which were mechanical and 440 biological), and this complication corresponded to 35% of all causes of prosthesis failure. Past medical history of mitral regurgitation, infective endocarditis, acute myocardial infarction, heavily calcified aortic annulus, and Marfan syndrome were the risk factors associated with a higher prevalence of leak2.

Given the importance of prosthetic valve leak among the complications of heart valve surgeries, it is hypothesized that this is a significant predictor of reoperation. The degree of leak seems to be implicated in early reoperation. However, no studies correlating the degree of prosthetic valve leak with possible postoperative events such as reoperation, infection or death are available.

Objective

To compare the postoperative outcome of patients with severe or mild/moderate prosthetic valve leak, and to verify whether differences in the clinical outcome of these patients exist in relation to the severity of the prosthetic valve leak.
Methods

In this retrospective study we analyzed the medical records from the database of the Clinical Unit of Heart Valve Disease of our institution, and other clinical data available in its electronic system. The following data were not available in these sources, and were therefore not analyzed: a) surgeries or clinical outcome of patients treated outside the institution; b) test results not available in the paper medical record that were being processed for inclusion in the electronic system by the time of data collection.

Inclusion criteria

• Patients undergoing heart valve surgery between 1999 and 2001 (called “Surgery 1”) who presented at least mild prosthetic valve leak (central and/or paravalvular), whether in biological or mechanical prosthesis;
• Surgery 1 was not necessarily the first heart valve surgery the patients had undergone in their lives;
• Leak may have occurred in the postoperative period of Surgery 1 or this surgery may have been performed precisely to correct a preexisting prosthetic valve leak.

Therefore, three possible situations could have occurred for each patient, as shown in Figure 1.

Exclusion criteria

Patients, who, despite meeting the inclusion criteria, had medical records with missing data that would affect the characterization of their clinical outcome or data analysis, were excluded from the study.

The leak degree was determined by pulsed and color Doppler according to the usual recommendations of the American Society of Echocardiography.

Patient groups

Patients who met the inclusion criteria and had medical records with complete data were distributed into two groups, according to the degree of prosthetic valve leak:
• Mi/Mo GROUP: patients presenting mild or moderate prosthetic valve leak.
• Sev GROUP: patients presenting severe prosthetic valve leak.

For group distribution purposes, the highest leak degree throughout the patient’s clinical / echocardiographic history was considered.

Patient selection

The patient selection process is summarized in Figure 2. A total of 1,350 medical records from patients undergoing heart valve surgery in our institution in the period between 1999 and 2001 were selected. Data from 185 patients who met the inclusion criteria of the protocol were evaluated. In accordance with the exclusion criteria, a total of 58 patients were enrolled in the study, 36 of whom in the Mi/Mo group and 22 in the Sev group.

Variables analyzed

Demographic, clinical, laboratory and echocardiographic variables were analyzed in both groups, namely:

Results

Of the 1,350 patients undergoing heart valve surgery in our institution between 1999 and 2001, 185 (13.7%) presented with prosthetic valve leak, which was graded from mild to severe. Most of the cases (86%) were paravalvular leaks. Considering the patients effectively selected for the study (n=58), the follow-up time was 4.49±2.98 years in the Mi/Mo group and 3.76±4.74 years in the Sev group (p=0.5167).

Mi/Mo and Sev group demographics are shown in Table 1. No statistically significant difference was found between the groups regarding age, gender, weight, or follow-up time in the study. In the preoperative period of Surgery 1, 31 patients from the Mi/Mo group had native valves, and five had prosthetic valves, whether with or without leak. In the Sev group, five patients had native valves and 17 had prosthetic valves, whether with or without leak (p<0.001, OR: 21.08 [95%CI 5.4 – 81.12] and RR: 5.56 [2.69 – 11.2]). This goes to show that the performance of surgical prosthetic valve replacement in patients already implanted with prostheses increases by 5.56 times the risk of significant prosthetic valve leak in the postoperative period.

Clinical and echocardiographic information of the patients selected are shown in Table 2. The patients did not differ as to the probability of death of any cause after Surgery 1 (p = 0.5238). We observed that patients with severe prosthetic valve leak had a 2.35-time higher
probability of reoperation than those with mild/moderate leak. However, the time for reoperation was surprisingly shorter (25.25±22.43 days) for the patients of the Mi/Mo group than for those of the Sev group (79.80±42.09 days) (p<0.001). In both groups, endocarditis was the major cause of reoperation, and biological aortic prostheses were those which more frequently presented leak, followed by biological mitral prostheses. Sixteen patients (72.7%) of the Sev group already presented prosthetic valve leak in the preoperative period of Surgery 1, in comparison with five patients (13.9%) of the Mi/Mo group. Therefore, we can observe that although the causes for the performance of Surgery 1 had been different, all patients who presented some prosthetic valve dysfunction in the Mi/Mo group also presented associated prosthetic valve leak, and most (16/17; 94.1%) of the patients of the Sev group with some prosthetic valve dysfunction also presented associated prosthetic valve leak. One patient in the Sev group did not present paravalvular or central prosthetic valve leak, despite undergoing surgical heart valve prosthesis replacement.

Table 2 also shows the mean number of heart valve surgeries performed in patients of the Mi/Mo and Sev groups (1.36±0.68 vs. 1.91±1.19 surgeries in the Mi/Mo and Sev groups, respectively; p=0.0551). We found no differences between the groups in relation to the number of heart valve surgeries performed throughout life (p=0.551), although a tendency to a greater number of surgeries among patients in the Sev group had been observed. When we classified the patients in relation to the performance of more than one heart valve surgery in life, we observed that nine patients (25%) of the Mi/Mo group had undergone more than one heart valve surgery in life, in comparison with 11 patients (50%) of the Sev group. A total of 34.5% of the patients in our study underwent more than one heart valve surgery.

The analysis of the Mi/Mo group showed a mean interval of 353±624 days between Surgery 1 and the diagnosis of prosthetic valve leak among the patients who presented leak after Surgery 1, whether or not presenting leak before this surgery. As regards the Sev group, the interval between Surgery 1 and the echocardiographic diagnosis of severe prosthetic leak was calculated at 230±433 days among patients in whom significant leak was absent in the preoperative period, but was

Table 1 – Profile of the patients selected

<table>
<thead>
<tr>
<th></th>
<th>Mi/Mo group (n=36)</th>
<th>Sev group (n=22)</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male gender</td>
<td>22 (61.1%)</td>
<td>15 (68.2%)</td>
<td>0.7932</td>
</tr>
<tr>
<td>Female gender</td>
<td>14 (38.9%)</td>
<td>7 (31.8%)</td>
<td></td>
</tr>
<tr>
<td>Age (years) by the time of Surgery 1</td>
<td>51.25±19.84</td>
<td>48.90±15.74</td>
<td>0.6204</td>
</tr>
<tr>
<td>Weight (kg)</td>
<td>75.7±20.4</td>
<td>68.2±15.2</td>
<td>0.267</td>
</tr>
<tr>
<td>Follow-up time (years) after Surgery 1</td>
<td>4.49±2.98</td>
<td>3.76±4.74</td>
<td>0.5167</td>
</tr>
<tr>
<td>Native valve pre Surgery 1</td>
<td>31</td>
<td>5</td>
<td>0.001*</td>
</tr>
<tr>
<td>Heart valve prosthesis pre Surgery 1</td>
<td>5</td>
<td>17</td>
<td>0.001*</td>
</tr>
</tbody>
</table>

*p in relation to the presence of heart valve prosthesis pre Surgery 1 (OR: 21.08 [5.4-81.12] and RR: 5.56 [2.69 – 11.2]).
Table 2 – Clinical and echocardiographic data of the patients selected

<table>
<thead>
<tr>
<th></th>
<th>Mi/Mo group (n=36)</th>
<th>Sev group (n=22)</th>
<th>Statistical analysis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Deaths of any cause after Surgery 1</td>
<td>3 (8.3%)</td>
<td>1 (4.5%)</td>
<td>OR=0.5238 (0.05-5.38)</td>
</tr>
<tr>
<td>Leak in the preoperative period of Surgery 1</td>
<td>5 (13.9%)</td>
<td>16 (72.7%)</td>
<td>OR=16.5333 (4.37-62.60)</td>
</tr>
<tr>
<td>Reoperation after Surgery 1</td>
<td>4 (11.1%)</td>
<td>5 (22.7%)</td>
<td>OR=2.3529 (0.56-9.94)</td>
</tr>
<tr>
<td>Time (days) from Surgery 1 to reoperation</td>
<td>25.25±22.43</td>
<td>79.80±42.09</td>
<td>p&lt;0.001</td>
</tr>
<tr>
<td>Cause of reoperation**</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Endocarditis</td>
<td>3 (75%)</td>
<td>3 (60%)</td>
<td></td>
</tr>
<tr>
<td>Bioprosthesis dysfunction and CHF*</td>
<td>1 (25%)</td>
<td>1 (20%)</td>
<td></td>
</tr>
<tr>
<td>Hemolysis**</td>
<td>-</td>
<td>1 (20%)</td>
<td></td>
</tr>
<tr>
<td>Total of heart valve surgeries pre and/or post-Surgery 1</td>
<td>1.36±0.68</td>
<td>1.91±1.19</td>
<td>p=0.0551</td>
</tr>
<tr>
<td>Prosthesis with leak*</td>
<td>24 AoBioP (55.8%)</td>
<td>15 AoBioP (57.7%)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>17 MiBioP (39.5%)</td>
<td>9 MiBioP (34.6%)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>1 AoMecP (2.3%)</td>
<td>1 AoMecP (3.8%)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>1 TriBioP (2.3%)</td>
<td>1 MiMecP (3.8%)</td>
<td></td>
</tr>
</tbody>
</table>

* Abbreviations: CHF = congestive heart failure; BioP = bioprosthesis; MecP = mechanical prosthesis; Ao = aortic; Mi = mitral; Tri = tricuspid.
** Values expressed as the number of patients with the clinical condition and percentage in relation to the total number of patients undergoing reoperation in each group. For example: of the four patients undergoing reoperation in the Mi/Mo group, three (75%) presented endocarditis; of the five patients undergoing reoperation in the Sev group, three (60%) presented endocarditis.
*** As diagnosed by increased lactic dehydrogenase and bilirubin levels and persistent hemoglobin reduction.

Discussion

The present study evaluated the profile of patients who presented severe prosthetic valve leak in relation to patients with mild or moderate leak in the postoperative period of heart valve surgery.

An OR = 16.5 (95% CI = 4.37-62.60) for the presence of leak as a preoperative diagnosis of Surgery 1 in the Sev group in relation to the Mi/Mo group confirmed that surgical prosthetic valve replacement is more frequent among patients with severe leak than among those with mild or moderate leak. Most of the patients (86.1%) with mild or moderate leak did not undergo Surgery 1 due to leak, but developed leak after this surgery, whether or not undergoing surgery later in time. Leak was the most frequent (72.7%) cause of Surgery 1 among patients with severe leak. Additionally, patients with severe leak have a greater chance of undergoing reoperation after surgical valve replacement, regardless of the reason for surgery (OR = 2.35; 95%CI = 0.56-9.94).

A statistically significant difference (p<0.001) was found for the time elapsed between Surgery 1 and reoperation in both groups. The Sev and Mi/Mo groups underwent reoperation 79.8±42.09 days and 25.25±22.43 days after Surgery 1, respectively. This is surprising, because we expected to find a shorter interval for reoperation in the Sev group. A hypothesis to explain this finding is that patients from the Sev group could be at higher surgical risk than those from the Mi/Mo group, which could occasionally postpone reoperation in the Sev group.
In both groups, more than 50% of the cases of prosthetic valve leak occurred in biological aortic prostheses, followed by biological mitral prostheses, in disagreement with Akins et al\textsuperscript{3} findings of a higher frequency of mitral leak (68%) in a series of 136 heart valve replacements. Also interesting is the fact that most of the cases that progressed with significant leak in the postoperative period of Surgery 1 presented prosthesis dysfunction, whether regurgitation alone or associated with stenosis (Table 1).

The comparison of the time elapsed between Surgery 1 and the diagnosis of prosthetic valve leak among patients who did not present leak at the first surgery showed a tendency to a shorter time for the diagnosis of severe leak in relation to less severe degrees of prosthetic valve leak (p=0.552), which is understandable in light of the greater severity and symptoms associated with severe prosthetic valve leak. Genoni et al\textsuperscript{3} demonstrated that among 598 patients, 22% of the cases of paravalvular leak were diagnosed in the first postoperative week, and 74% up to the first postoperative year. In our study, we verified that the mean time for the diagnosis of prosthetic valve leak in both groups was shorter than one year after surgery (353±624 days in the Mi/Mo group and 230±433 days in the Sev group).

Akin et al\textsuperscript{3} verified that 50% of the patients undergoing surgical correction of prosthetic valve leak had already undergone at least one cardiac surgery. In our study, we found 34.5% of the patients undergoing at least two heart valve surgeries in life. However, we should point out that our calculation is different from that of Akins et al\textsuperscript{3} for considering every surgery performed throughout the patients’ lives (and not only those performed before the surgical leak correction) and for being restricted to heart valve surgeries.

Complete resolution of leak after surgical heart valve replacement was less frequent among patients with severe leak (21.4%) than among patients with mild or moderate leak (40%) (OR = 0.5357, 95%CI = 0.07-4.2). However, there was a significant leak reduction by 72.7% in the cases with severe preoperative leak which progressed with some leak in the postoperative period of Surgery 1, naturally resulting in clinical improvement\textsuperscript{4}. Thus, as expected, the effectiveness of the surgical correction of prosthetic valve leak is higher in mild or moderate cases. Despite this fact, the percentage of patients progressing without prosthetic valve leak after Surgery 1 (40% in the Mi/Mo group and 21.4% in the Sev group; Table 4) was considerably lower than the values found by Akins et al\textsuperscript{3}. In this study, only 16% of the patients presented leak recurrence up to 10 years after surgical correction of the primary leak\textsuperscript{5}. Nevertheless, we should consider that the study did not include the cases of leak caused by endocarditis; also, it had a longer follow-up period (15 years)\textsuperscript{3}.

No consensus regarding the best way to treat prosthetic valve leak is available yet, especially in mild/moderate cases. Genoni et al study\textsuperscript{3} found mortality rates of 26% among patients who received conservative medical treatment and of 12% among those undergoing surgery. Akins et al\textsuperscript{3} concluded that the morbidity and mortality rates resulting from the surgical correction of prosthetic valve leak are acceptable. Some studies discuss the use of minimally invasive surgical techniques for heart valve replacement\textsuperscript{1-3} or improvements in the techniques using the open surgical approach in leak correction, especially for valve annulus reinforcement\textsuperscript{6}.

In addition to leak, other complications related to prosthetic valves contribute to the morbidity and mortality of patients undergoing surgical heart valve replacement. Of these, the most common are thromboembolism, hemorrhage, hemolysis, infectious endocarditis, heart valve failure and reoperation\textsuperscript{7-10}. Ten years after heart valve replacement, up to 50% of the patients require reoperation or die of complications related to the prosthetic heart valves. There is usually no difference between mechanical and biological prostheses, but the frequency and nature of the complications related to the prostheses vary according to type, model, and valve position, as well as to each patient’s characteristics\textsuperscript{4,11-15}.

By quantifying the comparative odds ratios between patients who presented varying degrees of prosthetic valve leak, the present study contributes to a better knowledge of its clinical outcome and provides information to guide therapies and establish prognoses. Although Genoni et al\textsuperscript{3} concluded that the surgical treatment of prosthetic valve leak seems to be better than medical treatment, the present study shows that the risk of reoperation and the effectiveness of surgical correction of heart valve replacement vary according to the leak degree. Furthermore, our study also contributed with the observation of the presence of an expressive number of cases of endocarditis as the cause of reoperation associated

### Table 3 – Laboratory data of Mi/Mo and Sev groups

<table>
<thead>
<tr>
<th></th>
<th>Mi/Mo group</th>
<th>Sev group</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hemoglobin (g/dL)</td>
<td>12.30±2.35</td>
<td>13.27±2.07</td>
<td>0.1063</td>
</tr>
<tr>
<td>Hematocrit (%)</td>
<td>36.86±6.33</td>
<td>39.45±5.29</td>
<td>0.1017</td>
</tr>
<tr>
<td>Creatinine (mg/dL)</td>
<td>1.1±0.42</td>
<td>1.25±1.21</td>
<td>0.5809</td>
</tr>
<tr>
<td>BUN (mg/dL)</td>
<td>50.09±23.34</td>
<td>53.89±45.17</td>
<td>0.7170</td>
</tr>
<tr>
<td>Glucose (mg/dL)</td>
<td>102.30±20.63</td>
<td>112.16±30.77</td>
<td>0.1968</td>
</tr>
</tbody>
</table>

### Table 4 – Effectiveness of surgical correction of prosthetic valve leak

<table>
<thead>
<tr>
<th></th>
<th>Mi/Mo group</th>
<th>Sev group</th>
<th>OR (95%CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of patients that already presented leak at Surgery 1</td>
<td>5/36 (13.9%)</td>
<td>16/22 (72.7%)</td>
<td>16.53 (4.37-62.60)</td>
</tr>
<tr>
<td>No leak in the postoperative period</td>
<td>2/5 (40%)</td>
<td>3/14 (21.4%)</td>
<td>2.44 (0.27-22.1)</td>
</tr>
<tr>
<td>Mi/Mo leak in the postoperative period</td>
<td>3/5 (60%)</td>
<td>8/14 (57.1%)</td>
<td>1.13 (0.14-9.00)</td>
</tr>
<tr>
<td>Sev leak in the postoperative period</td>
<td>0</td>
<td>3/14 (21.4%)</td>
<td></td>
</tr>
</tbody>
</table>

*Complete data on the postoperative outcome of 14/22 (63.6%)
both with the mild/moderate leak group (75%) and with the severe leak group (60%). Therefore, we point out that it is fundamental to consider the possibility of endocarditis in patients with postoperative prosthetic valve leak.\(^\text{16}\)

Our study was limited by the number of patients enrolled in each of the groups (n=36 and 22 in the Mi/Mo and Sev groups, respectively), so that statistically significant p values or odds ratios (OR) could not be obtained for some of the variables analyzed. This was due to the fact that prosthetic valve leak is relatively uncommon in our institution. However, with the values found we can point to tendencies and suggest further studies on this important topic. Additionally, it is important to recognize that a longer follow-up period may be necessary\(^\text{4}\) to compare the morbidity and mortality between the groups, since we analyzed approximately 4.49±2.96 years of the Mi/Mo group and 3.76±4.74 years of the Sev group. Other studies demonstrated a mean survival of only 30% 10 years after surgical correction of prosthetic valve leak.\(^\text{3}\)

**Conclusions**

(1) Surgical correction of prosthetic valve leak is more frequent in patients with severe leak than in those with mild or moderate leak. The presence of prosthesis dysfunction is a predisposing factor for the development of severe postoperative leak.

(2) Patients who already had or who will develop severe prosthetic valve leak have a higher chance of undergoing reoperation after surgical heart valve replacement, regardless of the reason for surgery.

(3) Severe prosthetic valve leak is more difficult to resolve completely after surgical treatment.

(4) Endocarditis was the most common cause of reoperation, regardless of the degree of prosthetic valve leak.

(5) A higher incidence of prosthetic valve leak was observed in aortic bioprostheses in the series of surgeries performed in our institution.

**Potential Conflict of Interest**

No potential conflict of interest relevant to this article was reported.

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There were no external funding sources for this study.

**Study Association**

This study is not associated with any post-graduation program.

**References**


