

Systematic review and meta-analysis of surgical outcomes in Marfan patients undergoing aortic root surgery by composite-valve graft or valve sparing root replacement

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Background: A major, life-limiting feature of Marfan syndrome (MFS) is the presence of aneurysmal disease. Cardiovascular intervention has dramatically improved the life expectancy of Marfan patients. Traditionally, the management of aortic root disease has been undertaken with composite-valve graft replacing the aortic valve and proximal aorta; more recently, valve sparing procedures have been developed to avoid the need for anticoagulation. This meta-analysis assesses the important surgical outcomes of the two surgical techniques.

Methods: A systematic review and meta-analysis of 23 studies reporting the outcomes of aortic root surgery in Marfan patients with data extracted for outcomes of early and late mortality, thromboembolic events, late bleeding complications and surgical reintervention rates.

Results: The outcomes of 2,976 Marfan patients undergoing aortic root surgery were analysed, 1,624 patients were treated with composite valve graft (CVG) and 1,352 patients were treated with valve sparing root replacement (VSRR). When compared against CVG, VSRR was associated with reduced risk of thromboembolism (OR =0.32; 95% CI, 0.16–0.62, P=0.0008), late hemorrhagic complications (OR =0.18; 95% CI, 0.07–0.45; P=0.0003) and endocarditis (OR =0.27; 95% CI, 0.10–0.68; P=0.006). Importantly there was no significant difference in reintervention rates between VSRR and CVG (OR =0.89; 95% CI, 0.35–2.24; P=0.80).

Conclusions: There is an increasing body of evidence that VSRR can be reliably performed in Marfan patients, resulting in a durable repair with no increased risk of re-operation compared to CVG, thus avoiding the need for systemic anticoagulation in selected patients.

Keywords: Marfan syndrome (MFS); valve sparing root; David procedure; Bentall; composite valve graft (CVG); total root replacement



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Introduction

Marfan syndrome (MFS) is a genetic disorder of connective tissues with autosomal dominant inheritance due to mutation of the FBN1 gene, affecting approximately 1 in 5,000 patients. Diagnosis of the condition is made using the modified Ghent-2 criteria which provides a diagnosis based on clinical features, family history and genetic testing (1). Historically, patients with MFS had a median life expectancy of 49 years, limited largely due to complications of aortic pathology. Aneurysmal disease of the ascending aorta is particularly common with ascending aortic dilatation present in 53% of patients at 30 years. Life expectancy for Marfan patients is now similar to that of the general population, due to appropriate cardiovascular surgical intervention to minimize the risk of aortic complications (2).

Elective aortic root replacement is indicated in patients with MFS with an aortic root aneurysm ≥ 50 mm in diameter, or ≥ 45 mm when the patient has significant risk factors such as family history of acute aortic syndrome or rapid aneurysmal expansion of >3 mm/year (3). Management of aortic root pathology necessitates resection of diseased aortic tissue, with the historical gold standard being total root replacement with a modified Bentall procedure as originally described in 1968 by Bentall and De Bono (4). This procedure replaces the aortic valve and proximal aorta with a composite valve graft (CVG), to which the coronary arteries are re-implanted. Due to the young age of Marfan patients, the use of a mechanical valve is often necessary, placing the patient at risk of thromboembolic and hemorrhagic complications. More recently, techniques to preserve the aortic valve for individuals with isolated aneurysmal disease and normal or mildly abnormal aortic valve leaflets have been developed (5,6). These methods obviate the need for anticoagulation and avoid the risk of valve-related thromboembolism. However, a major concern of preserving aortic valve leaflets known to contain defective fibrillin-1 in MFS patients is the risk of early repair failure and need for re-operation (7). The concerns regarding the durability of valve sparing root replacement (VSRR) are being addressed with increased experience in the techniques and longer-term follow up data. Given the overall small number of MFS patients undergoing aortic root surgery, meta-analysis of the outcomes of observational studies provides the best evidence to support decision-making in this group of patients. To date, two

meta-analyses have been undertaken by Benedetto *et al.* (8) in 2011 and Hu *et al.* (9) in 2014. This systematic review and meta-analysis aims to include an update of current published data and outcomes.

Methods

Literature search strategy

This meta-analysis was performed in accordance with PRISMA recommendations and guidance. The search strategy was employed to search electronic databases EMBASE, Ovid Medline, the entire Cochrane Central Register of Controlled Trials (CCRCT), Cochrane Database of Systematic reviews (CDSR), the Database of Abstracts of Reviews of Effects (DARE) and the ACP journal club from their inception to September 2017. The search strategy included search terms for ((aortic valve sparing or aortic valve preserving or David procedure) OR (Bentall or aortic root replacement or CVG) OR (aortic root surgery)) AND Marfan. The bibliography of previous systematic reviews was assessed to ensure no additional publications are missed.

Selection criteria

Eligibility for inclusion in this systematic review and meta-analysis included papers that assessed the outcomes of patients undergoing aortic root surgery in adult patients by either VSRR of any technique or total root replacement with CVG. In order to ensure sufficient institutional experience, papers were only included if more than 30 cases were reported. Studies were excluded if there was inadequate data regarding the outcomes of the repair technique or there was no separation of Marfan patient outcomes from the general patient outcomes. When centres reported outcomes of overlapping patient series then the most contemporary series was analysed. The analysis was limited to English language papers. Conference abstracts, case reports, editorials, expert opinion, reviews and expert opinion were excluded.

Data extraction

For the assessed papers, data was extracted from the reviewed text, tables and figures. Data was extracted independently by two of the authors (CDF and AWS) and any discrepancies were reviewed and discussed until consensus was reached.

The recorded parameters were: number of cases in series, procedure undertaken, urgency of procedure, average age, average follow up, early death, late death, bleeding complications (late), systemic thromboembolic complications, post-operative endocarditis and reoperation.

Statistical analysis

Meta-analysis of incidence rates of post-operative complications, including endocarditis, thromboembolism, hemorrhagic complication and reoperation, was performed. The incidence rates were reported as number of events/follow up year. Incidence rates were assessed drawing data from all papers. Incidence data was assessed using Comprehensive Meta-analysis v3.3. Comparative outcome data was determined only from observational studies reporting outcomes of both VSRR and CVG in MFS patients. Comparative outcomes were reported in events per follow-up year and were assessed using Review Manager v5.3 (Cochrane Collaboration, Copenhagen, Denmark). Data heterogeneity was assessed with the Cochrane Q statistic with P value <0.05 being significant and the I^2 test statistic. Publication bias was assessed by generation of funnel plots and assessment using Egger's test. The number of patient years was calculated by multiplying the number of patients included in the study by the average follow up duration of that study. Annual event rates are determined from data on all included studies, with no comparison made between CVG and VSRR on overall event rate data due to the risk of heterogeneity of data. Comparative outcomes are determined from subgroup random-effects analysis of the included studies reporting outcomes of both CVG and VSRR to determine the effect of surgical technique on outcomes. Sensitivity analysis was conducted for all assessed variables using 'remove-one' analysis. Individual patient survival data was reconstructed using an iterative algorithm that was applied to digitized source Kaplan-Meier curves and subsequently aggregated and graphed (10).

Results

The search strategy revealed 900 citations for review after duplicates were removed, 23 of which met the pre-determined inclusion criteria (11-33) (*Figure S1*). The included publications had a total patient population of 2,976; 1,624 patients were treated with CVG and 1,352 patients were treated with VSRR. For all patients, the total number of patient-years follow up for CVG is

8,794 and the total number of patient-years follow up for VSRR is 5,741. The average follow-up for period for CVG in 13 studies reporting follow up duration was 7.14 years and the average follow up period for VSRR in 20 studies reporting follow up duration was 4.5 years. There were 2 studies reporting the outcomes of CVG only (11,19) and 8 studies reporting the outcomes of VSRR only (15,17,20,22,23,26,28,31). Thirteen studies reported outcomes for both CVG and VSRR (12-14,16,18,21,24,25,27,29,30,32,33). The overall study characteristics are detailed in *Table 1*.

Assessing overall data, the incidence of late bleeding complications for CVG was 1.3% (95% CI, 0.4–2.1%) events/follow up year and the incidence of thromboembolic phenomena was 0.7% (95% CI, 0.5–0.9%) events/follow up year. The incidence of late bleeding complications and thromboembolic phenomena for VSRR was 0.1% (95% CI, 0.0–0.3%) and 0.4% (95% CI, 0.3–0.5%) event/follow-up year respectively. The incidence of re-operation for CVG and VSRR was 1.3% (95% CI, –1.0–3.7%) and 0.6% (95% CI, 0.3–0.9%) events/follow up year respectively. Full event rates for surgical outcomes are detailed in *Table 2*.

Meta-analysis of comparative studies demonstrated a significant reduction in late mortality for patients undergoing VSRR (OR =0.26; 95% CI, 0.15–0.46; $P<0.0001$) (*Figure 1*). As expected, there was a reduced risk of thromboembolic events (*Figure 2*) for VSRR (OR =0.32; 95% CI, 0.16–0.62; $P=0.0008$) and late hemorrhagic complications (*Figure 3*) (OR =0.18; 95% CI, 0.07–0.45; $P=0.0003$). The odds of post-operative endocarditis were less after VSRR than after CVG (*Figure 4*) (OR =0.27; 95% CI, 0.10–0.68; $P=0.006$). Importantly, the risk of re-operation was not increased (*Figure 5*) for VSRR when compared to CVG (OR =0.89; 95% CI, 0.35–2.24; $P=0.80$), however this was a high level of heterogeneity between studies ($I^2=60\%$), sensitivity analysis did not reveal any change in combined effect. Follow-up duration of VSRR and CVG did not explain any difference between VSRR and CVG on meta-regression analysis ($R^2=0.00$).

The congregate Kaplan-Meier curve for overall survival for CVG and VSRR is shown in *Figure 6*. Overall survival at 1-, 2-, 3-, 5-, and 10-year was 97.6%, 95.9%, 95.1%, 92.4%, and 84.5%, respectively. Congregate Kaplan-Meier curves for overall re-intervention rate for CVG and VSRR (*Figure 7*) was 97.8%, 97.0%, 96.2%, 95.6%, and 90.5%, respectively at 1-, 2-, 3-, 5-, and 10-year.

There was no evidence of publication bias in any of the reported outcomes using Egger's test or through visual

Table 1 Characteristics of studies reporting outcomes of Mafan patients undergoing aortic root surgery with composite valve graft or valve sparing root replacement

First Author	Year	Study location	Sample Size	Study period	Average follow-up (years)	Average follow-up CVG (years)	Average follow-up VSRR (years)	CVG procedures total	% emergency/urgent	VSRR procedures total	% emergency/urgent	Average patient age CVG	Average patient age VSRR
Alexiou	2001	Southampton, UK	65	1972–1998	8	8	NA	65	41.5	0	NA	41.7	NA
Bernhardt	2011	Hamburg, Germany	88	1998–2010	3.2	NA	NA	30	53.3	58	17.2	40.8	37.7
Cameron	2009	Maryland, USA	372	1976–2006	NA	NA	NA	287	NA	85	0	NA	NA
Coselli	2014	Houston, USA	316	2005–2010	1	1	1	77	23.4	239	4.2	39.0	33.0
David	2015	Toronto, Canada	149	1988–2012	10	NA	10	0	NA	149	0	NA	35.7
De Oliveira	2003	Toronto, Canada	44	1988–2001	NA	6.25	4.1	44	36.4	61	0	34.0	NA
Forteza	2010	Madrid, Spain	37	2004–2009	2.25	NA	2.25	0	NA	37	0	NA	30.0
Gott	1999	Baltimore, USA	675	1968–1997	6.7	6.7	NA	625	NA	50	NA	34.0	34.0
Groenik	1999	Amsterdam, Netherlands	125	1986–1996	8.9	8.9	NA	43	30.2	0	NA	30.0	NA
Kallenbach	2007	Heidelberg, Germany	59	1993–2005	4.5	NA	4.5	0	NA	59	0	NA	30.0
Karck	2004	Hannover, Germany	119	1979–2002	NA	9.5	2.5	74	23.0	45	6.7	28.0	35.7
Kari	2014	Freiburg, Germany	48	1997–2013	3.8	NA	3.8	0	NA	48	12.5	NA	33.0
Kunihara	2012	Homburg, Germany	33	1995–2011	5.5	NA	5.5	0	NA	33	24.2	NA	31.0
Nardi	2010	Rome, Italy	47	1998–2008	NA	6.25	4.5	23	17.4	24	12.5	38.0	36.0
Price	2016	Baltimore, USA	165	1997–2014	7.8	7.1	8.4	67	23.9	98	3.1	37.0	36.0
Schmidtke	2012	Lubeck, Germany	37	1993–2009	3.5	NA	3.5	0	NA	37	32.4	NA	32.0
Schoenhoff	2015	Bern, Switzerland	59	1995–2014	NA	8.8	6.5	30	0	29	0	36.0	27.0
Settepani	2007	Rome, Italy	35	2000–2006	1.6	NA	1.6	0	NA	35	5.7	NA	36.5
Shimizu	2012	Tokyo, Japan	50	1987–2010	NA	NA	NA	37	NA	13	NA	NA	NA
Song	2014	Oregon, USA	288	1981–2012	10.5	10.5	6.2	113	31.9	87	4.6	36.3	31.0
Urbanski	2017	Bad Neustadt, Germany	42	2002–2016	6.1	NA	6.1	0	NA	42	9.5	NA	43.0
Volguina	2009	Stanford, USA	151	2005–2008	0.08	0.08	0.08	46	23.9	105	5.7	39.0	31.0
Zehr	2005	Rochester, USA	83	1971–2001	7.6	7.6	7.6	63	NA	18	NA	NA	NA

CVG, composite valve graft; VSRR, valve sparing root replacement; NA, not available/appropriate.

Table 2 Summary table of clinical outcomes of all patients included in the present meta-analysis

First author	Year	CVG early mortality	VSRR early mortality	CVG late mortality	VSRR late mortality	CVG reoperation	VSRR reoperation	CVG late bleeding event	VSRR late bleeding event	CVG thrombo-embolism	VSRR thrombo-embolism	CVG endocarditis	VSRR endocarditis
Alexiou	2001	0	–	15	–	2	–	5	–	5	–	1	–
Bernhardt	2011	0	0	3	2	6	2	NA	NA	2	0	3	0
Cameron	2009	2	0	72	2	11	4	NA	NA	19	0	18	0
Coselli	2014	1	1	4	2	0	1	5	3	2	4	1	0
David	2015	–	1	–	6	–	1	–	NA	–	6	–	1
De Oliveira	2003	0	0	5	2	5	1	NA	NA	4	0	3	0
Forteza	2010	–	0	–	1	–	0	–	0	–	0	–	0
Gott	1999	22	0	110	4	12	1	2	0	26	1	24	0
Groenik	1999	5	–	3	–	1	–	NA	–	5	–	NA	–
Kallenbach	2007	–	0	–	5	–	7	–	4	–	0	–	0
Karck	2004	5	NA	9	3	7	4	12	0	5	1	2	NA
Kari	2014	0	1	–	3	–	1	–	NA	–	NA	–	NA
Kunihara	2012	0	0	–	0	–	3	–	0	–	1	–	0
Nardi	2010	1	0	2	0	0	2	NA	NA	0	0	0	1
Price	2016	0	0	NA	NR	4	4	7	0	8	4	2	1
Schmidtke	2012	–	0	–	0	–	3	–	0	–	2	–	0
Schoenhoff	2015	1	0	3	1	0	4	2	0	2	0	0	0
Settepani	2007	–	0	–	0	–	3	–	NA	–	0	–	1
Shimizu	2012	NA	0	NA	NA	NA	NA	3	0	2	0	NA	NA
Song	2014	NA	NA	NA	NA	13	0	NA	NA	NA	NA	NA	NA
Urbanski	2017	–	0	–	0	–	2	–	0	–	0	–	0
Volguina	2009	0	0	NA	NA	0	0	NA	NA	0	1	0	0
Zehr	2005	NA	NA	NA	NA	2	5	12	0	5	0	1	0
No. studies reporting outcome		11	18	9	16	14	20	8	10	13	19	12	17
Total number of patients		1,124	1,234	1,014	1,031	1,300	1,299	1,038	700	1,511	1,217	1,144	1,159
Annual event rate (95% CI)		0.003 (0.002–0.005)	0.001 (0.000–0.002)	0.018 (0.011–0.025)	0.005 (0.003–0.007)	0.013 (–0.01–0.037)	0.006 (0.003–0.009)	0.013 (0.004–0.021)	0.001 (0.000–0.003)	0.007 (0.005–0.009)	0.004 (0.003–0.005)	0.004 (0.003–0.005)	0.001 (0.000–0.002)
CVG, composite valve graft; VSRR, valve sparing root replacement; NA, not available/appropriate; CI, confidence interval.													

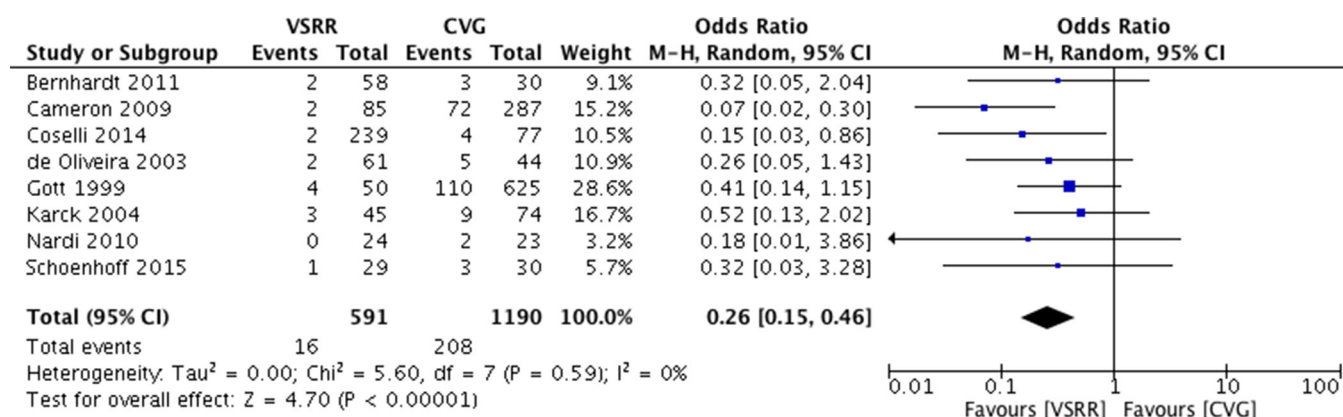


Figure 1 Comparison of late mortality between CVG and VSRR. CVG, composite valve graft; VSRR, valve sparing root replacement.

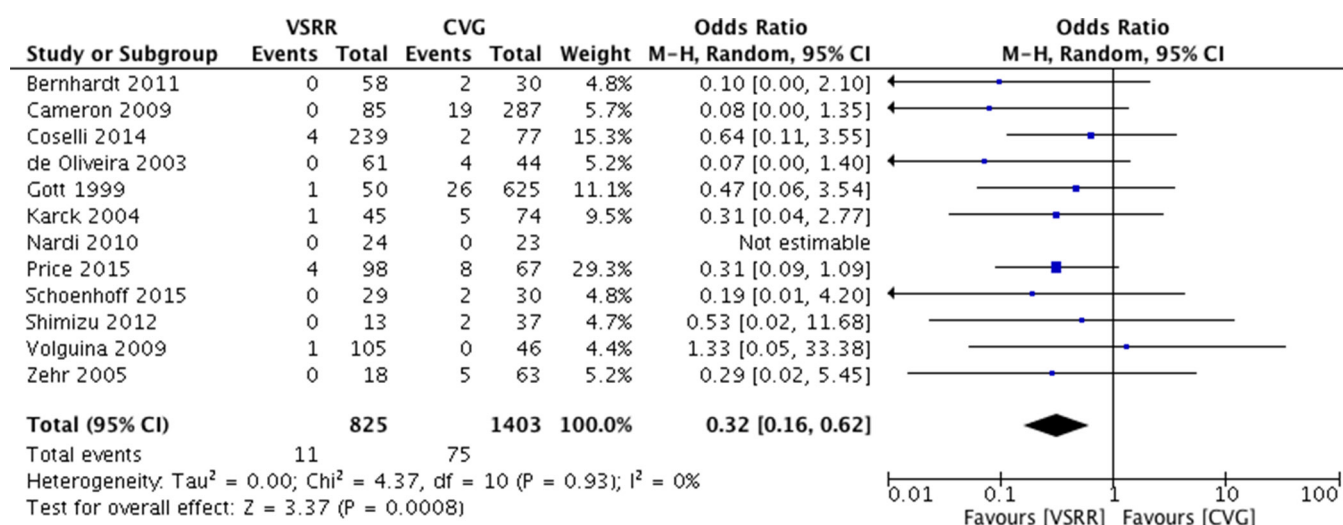


Figure 2 Comparison of thromboembolic events between CVG and VSRR. CVG, composite valve graft; VSRR, valve sparing root replacement.

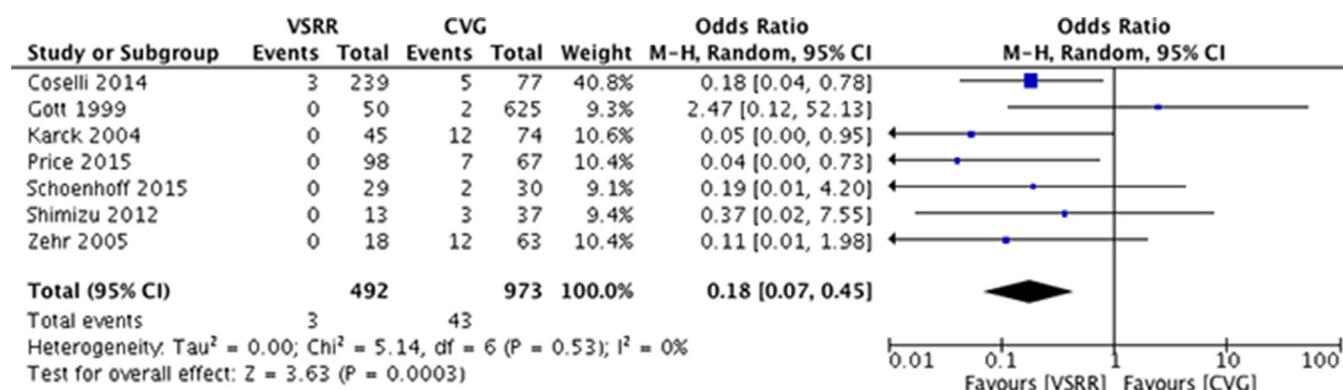


Figure 3 Comparison of late haemorrhagic events between CVG and VSRR. CVG, composite valve graft; VSRR, valve sparing root replacement.

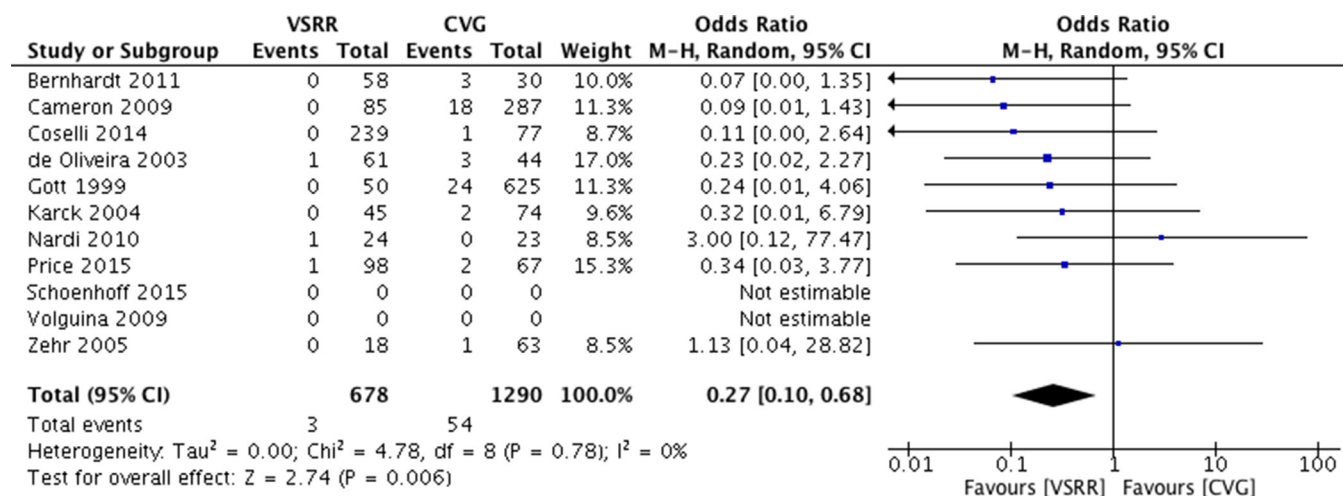


Figure 4 Comparison of endocarditis events between CVG and VSRR. CVG, composite valve graft; VSRR, valve sparing root replacement.

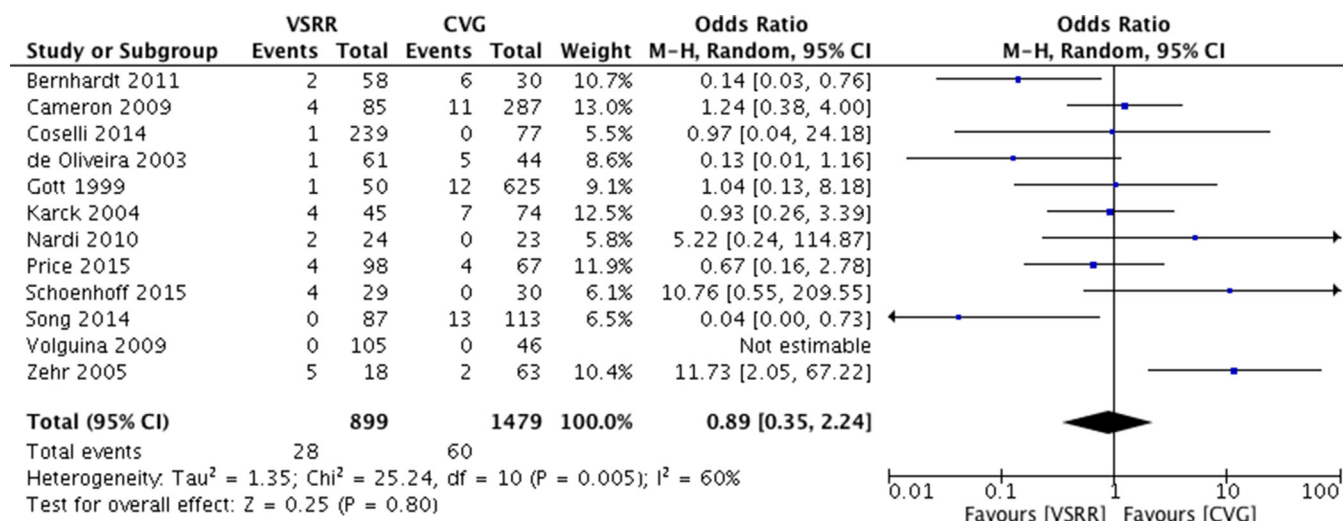


Figure 5 Comparison of surgical re-intervention between CVG and VSRR. CVG, composite valve graft; VSRR, valve sparing root replacement.

inspection of funnel plots (Figures S2-S7).

Discussion

Vascular intervention in patients with MFS is responsible for the recent and substantial improvement in the life-expectancy of this young population. For a condition affecting a small proportion of the population, the prospect of a high quality randomized control trial to provide evidence is unlikely. However, by pooling data from observational studies for meta-analysis, we are able to

improve the evidence-base upon which we make decisions for intervention strategy. The present study has drawn data from a pool of 2,976 patients from experienced surgical centers to determine incidence rates for important post-operative complications and data from 2,457 patients in studies that provide a comparison of outcomes of composite valve-graft and valve-sparing root replacement within the same institution.

This meta-analysis has demonstrated the potential long-term benefit of Marfan patients undergoing valve-sparing root replacement over composite valve grafts. Significant

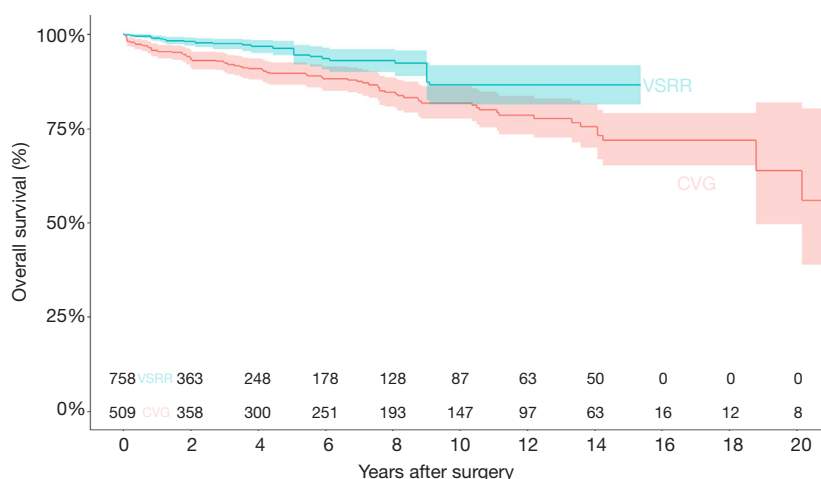


Figure 6 Congregate Kaplan-Meier curve for overall survival between CVG and VSRR. Shaded region represents 95% confidence interval. CVG, composite valve graft; VSRR, valve sparing root replacement.

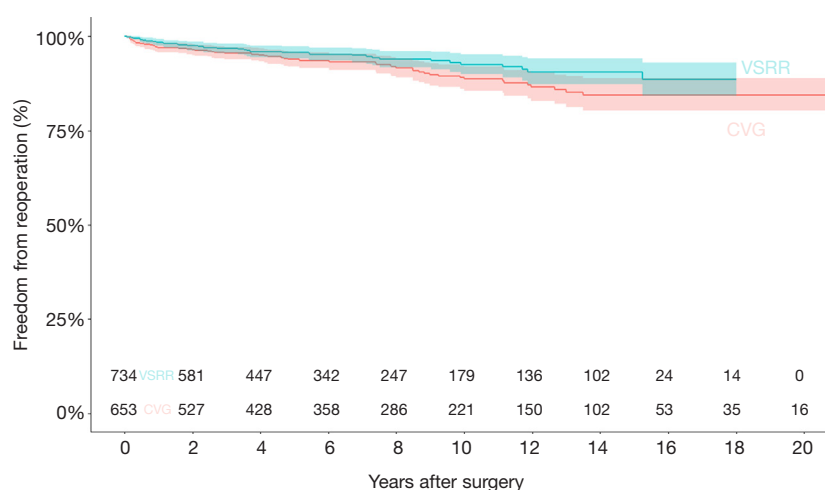


Figure 7 Congregate Kaplan-Meier curve for late re-operation for VSRR and CVG. Shaded region represents 95% confidence interval. VSRR, valve sparing root replacement; CVG, composite valve graft.

benefit of VSRR for thromboembolic complications have previously been demonstrated in meta-analysis (8,9), the latter study by Hu *et al.* in 2014 (9) also favoured VSRR for endocarditis rates. Interestingly, Benedetto *et al.* (8) reported a four-fold increased risk of intervention on aortic root with VSRR compared to CVG, a finding not supported by the present meta-analysis or by Hu *et al.* This disparity may be explained by the greater sample sizes available in the more recent studies.

The traditional gold standard of composite valve graft for the management of aortic root disease provides an

excellent improvement in the survival of Marfan patients. However, due to their young age at the time of surgery, they usually require the use of a mechanical prosthesis and systemic anticoagulation which puts the individual at risk of hemorrhagic and thromboembolic complications as well as an increased risk of endocarditis. Furthermore, female patients may be of childbearing age, making the provision of systemic anticoagulation challenging.

Limitations in current data stem from the observational and largely retrospective nature of the data analyzed which increases the risk of bias and potentially decreases the

accuracy of collected information. Volguina and Coselli reported early outcomes of a multicenter, prospectively designed study of aortic surgery in Marfan patients from centers in North America, South America and Europe (14,32). This study along with the GenTAC registry (34), a prospectively designed multicenter registry for genetically mediated thoracic aneurysmal disease will certainly add to the quality of evidence to guide decision making when long-term data becomes available.

An important concession for the decision-making process is that valve-sparing root replacement is by no means appropriate for every Marfan patient undergoing aortic surgery as both surgeon and patient factors may necessitate a total root replacement with composite valve-graft. Not all patients are appropriate for valve-sparing root replacement as the presence of leaflet fenestration or tissue redundancy may make a satisfactory and durable repair impossible. This inherent selection bias will be impossible to eliminate from any future study. Furthermore, because valve sparing surgery is a technically demanding procedure, centers with low volumes are likely to have worse outcomes (35). Therefore the procedure should be carried out in centers with sufficient expertise.

Expert opinion 1 (Tirone E. David)

This is a timely meta-analysis on the surgical outcomes of patients with MFS undergoing aortic root surgery by composite-valve graft (CVG) or VSRR. Flynn and colleagues reviewed 900 references and identified 23 studies that they used to compare the outcomes of 1,624 patients with MFS treated with CVG with 1,352 patients who had VSRR. The average follow-up was longer for CVG (7.1 years) than for VSRR (4.5 years) patients. As expected, rates of bleeding and thromboembolism were higher in CVG than in VSRR patients. Unexpectedly, the rate of reoperation was also higher in CVG than in VSRR, but as pointed out by the authors there was a high level of heterogeneity between studies. I would have expected a higher rate of reoperation after VSRR.

I believe that patients with aortic root aneurysm associated with genetic syndromes should have VSRR whenever possible. Actually, it is feasible in most patients because surgery is recommended early in the course of aortic sinuses dilatation when the cusps are still normal or have minor structural abnormalities (36). I have, and continue to be very conservative in all patients with aortic root aneurysm and perform VSRR only when the cusps are

normal or have only minor abnormalities such as elongation of the free margin or small stress fenestrations along the commissural areas (37). This conservative approach may explain my exceptionally good long-term results as they relate to reoperation and the development of aortic insufficiency (36,37).

CVG was introduced two decades earlier than VSRR and it is a more reproducible operation than VSRR. The main drawback of CVG is that mechanical valves should be used in patients with MFS because they are often in their second or third decades of life by the time they need aortic root surgery and mechanical valves require lifelong anticoagulation with warfarin. Tissue valves in these patients have limited durability.

Expert opinion 2: “Don’t throw the baby out with the bathwater” (George Matalanis)

The general philosophy of VSRR is the provision of a hemodynamically efficient aortic valve with durability exceeding that of a bioprosthesis, no requirement for anticoagulation and low risk of long term complications such as infection. Because most Marfan’s patients present for surgery at a young age, the avoidance of anticoagulation is not only desirable for considerations such as life style and pregnancy, but also from the inevitable need for subsequent cardiac and non-cardiac surgery. The greater exercise demands in this patient group also benefit from the absence of significant gradients or regurgitation from a well performed VSRR. Nonetheless, in the context of the well described Marfan’s fibrillin deficiency and frequent occurrence of fenestrations there is a genuine concern regarding durability.

In this meta-analysis, Flynn and colleagues performed a systematic review of the available literature to elucidate both the acute and chronic performance of VSRR *vs.* CVG. Importantly, the operative mortality was very low and equivalent in both groups, as expected from this young and healthy cohort. Thus, the extra manipulations incurred in performing a VSRR were not resulting in extra operative risk. The rest of the findings supported better outcomes with VSRR including, lower long-term bleeding, thromboembolic and infective complications than CVG. All this was achieved without a significant trade off of increased late re-operation rate.

In the midst of such salutary findings in favour of valve preservation in the Marfan’s root, a word of caution is in order. Firstly, it is sobering to appreciate that such

great long-term outcomes of VSRR would have easily been lost had the early morbidity and mortality not been so excellent in the reported series. Such results are achieved in experienced centers due to a combination of careful case selection and surgical familiarity with the techniques. Clearly, there are Marfan's aortic valves with numerous fenestrations and extreme fragility, that would be inappropriate to preserve. Equally obvious, as pointed out by the authors, there was a degree of variability in the freedom for re-intervention between the series. Therefore, we will have to wait for larger numbers followed for longer periods to be absolutely sure of the longevity of VSRR in Marfan's syndrome.

The authors have laid down a firm evidence basis for us to recommend VSRR in centres with good experience in the repair techniques, with the same confidence as mitral valve repair in Marfan's syndrome.

Expert opinion 3: selection and expertise is the key to success (Martin Misfeld)

Complications of aneurysmal disease limits the life expectancy of patients with MFS. Replacement of the aortic root either by CVG replacement or by VSRR have led to excellent surgical results.

This systematic review and meta-analysis of patients with MFS was performed by Flynn *et al.* with a total number of 23 studies, incorporating 1,624 CVG and 1,352 VSRR patients. There was no statistically significant difference in re-intervention rates. There was a significantly reduced risk of thromboembolism, late hemorrhagic complications and endocarditis in patients with VSRR.

A classic phenotypic expression of MFS affects mutations of the gene encoding fibrillin-1. Recent genetic studies demonstrate a high degree of overlap between MFS other connective tissue diseases (e.g., Loeys-Dietz syndrome, Ehlers-Danlos syndrome, and others). There are several variations of clinical phenotype in patients with MFS, even intra-familial. Therefore, the diagnosis of MFS may not represent a uniform disease and the clinical picture and indication for surgery may also vary.

In this manuscript, VSRR has been shown to have advantages compared to CVG, although there is clearly a selection bias. VSRR itself contains the aortic valve remodelling and the reimplantation technique. Stabilization of the aortic root annulus is a key factor in MFS patients. Therefore, either the reimplantation technique or the remodelling technique with additional annular support

should be performed. Surgical expertise is essential when performing this procedure, particularly when repair to aortic valve cusps is required. Regular follow-up in qualified centers for surveillance of the residual aorta and dealing with concomitant diseases (e.g., skeletal and eyes) as well as genetic and family counseling is required.

As the authors rightly state, patients with MFS should be treated in centers with experience managing this complicated group of patients. This allows for appropriate decision making for surgery, enables additional support for the management of concomitant diseases and an adequate surgical expertise to perform either CVG or VSRR to obtain the excellent results presented in this paper.

Expert opinion 4 (Stefano Mastrobuoni and Gebrine El Khoury)

Dr. Flynn and colleagues have performed a systematic review and meta-analysis of studies on the surgical outcomes of VSRR versus traditional operation (Bentall operation) with valve prosthesis in patients with Marfan syndrome. The current meta-analysis is particularly important because since a previous analysis by Benedetto (8) in 2011 that questioned the efficacy of VSRR in this selective cohort of patients, several important papers have been published which have been included by Flynn and colleagues. Indeed, since then, Dr. David (15) published his experience with Marfan patients as well as the groups of Dr. Cameron (25) in Baltimore (US), Dr. Carrel (27) in Bern (Switzerland), Dr. Sievers (26) in Lubeck (Germany) and others. The current meta-analysis shows clearly that VSRR is associated with particularly low rates of valve-related complications such as bleeding, thromboembolism, endocarditis and an improved survival compared to the conventional Bentall operation. The relative risk reduction is stunningly in the order of 70% for these complications. It seems unlikely that future studies may overturn these results. Further, the current analysis did not reveal an increased risk of reoperation after VSRR that is usually considered a major limitation of this operation.

Usually Marfan patients arrive at surgery in their 3rd or 4th decade of life. In this cohort of young patients requiring root and ascending aorta replacement the choice is between a composite graft with either a mechanical or a tissue prosthesis, VSRR or a Ross operation. A composite graft with a tissue prosthesis at this age expose the patient to a high risk of reoperation in the mid-term and a decreased survival, and therefore offers no advantages compared to

VSRR. Similarly, considering the underlying connective tissue disorder and the complexity of the procedure, the Ross operation is not an attractive option either. Thus, the choice really is between VSRR and mechanical prosthesis. We believe that the reduction of valve-related complications associated with mechanical prosthesis and life-long oral anticoagulation is of particular interest in order to guarantee the quality of life of these young, active patients even if they may have a certain risk of reoperation over time. Nonetheless, in experienced centers, reoperations after VSRR are routinely performed with low morbidity and mortality. Moreover, we have observed that Marfan patients present for surgery early in the disease course and large fenestrations or other cusp diseases that preclude an effective repair are rarely seen. Therefore, in elective Marfan patients VSRR can be safely performed with excellent results.

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None.

Footnote

Conflicts of Interest: The authors have no conflicts of interest to declare.

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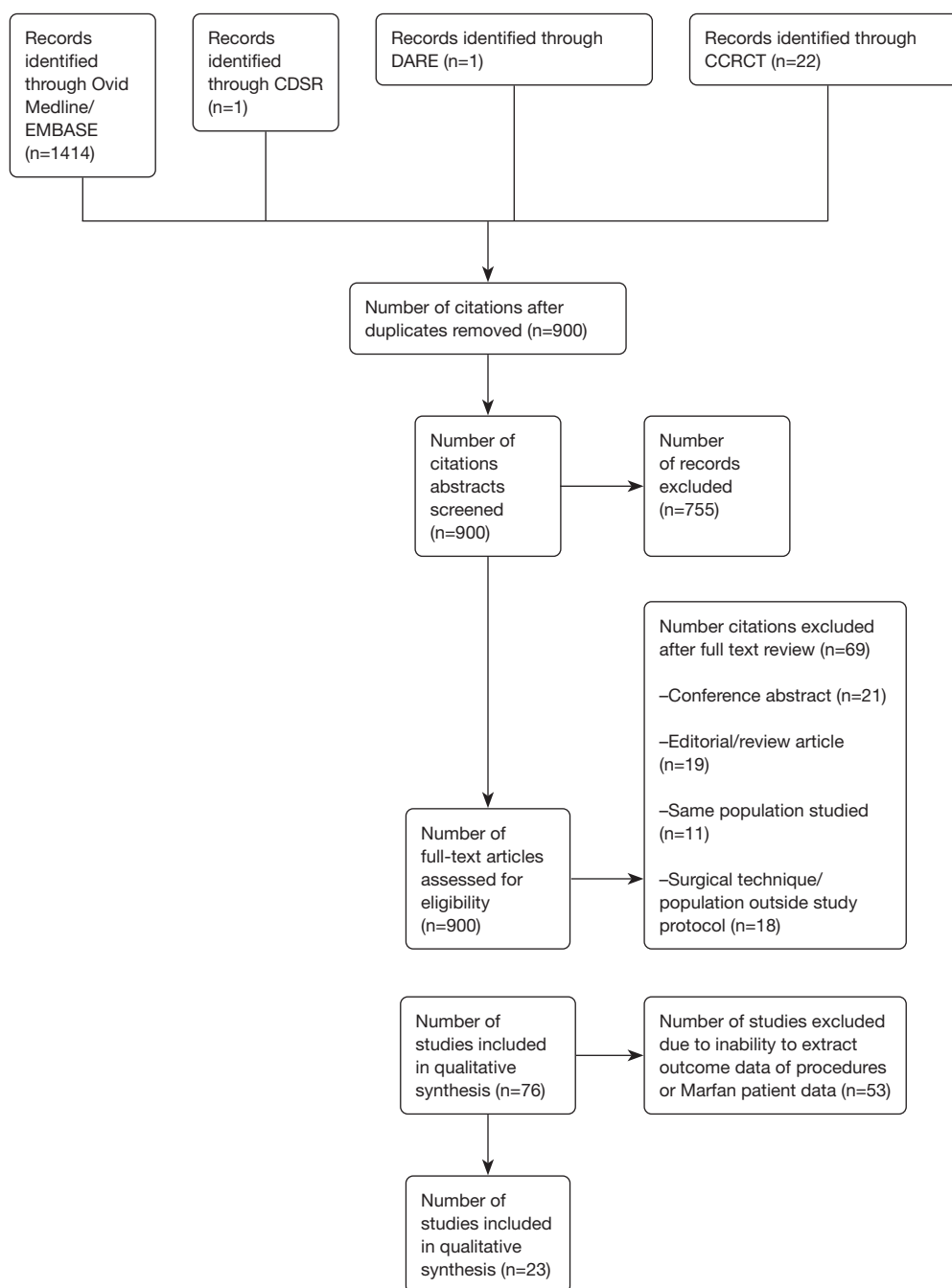


Figure S1 Study selection. Flow chart detailing the steps of systematic review to identify studies reporting the outcomes of composite valve graft and valve sparing root replacement in Marfan syndrome.

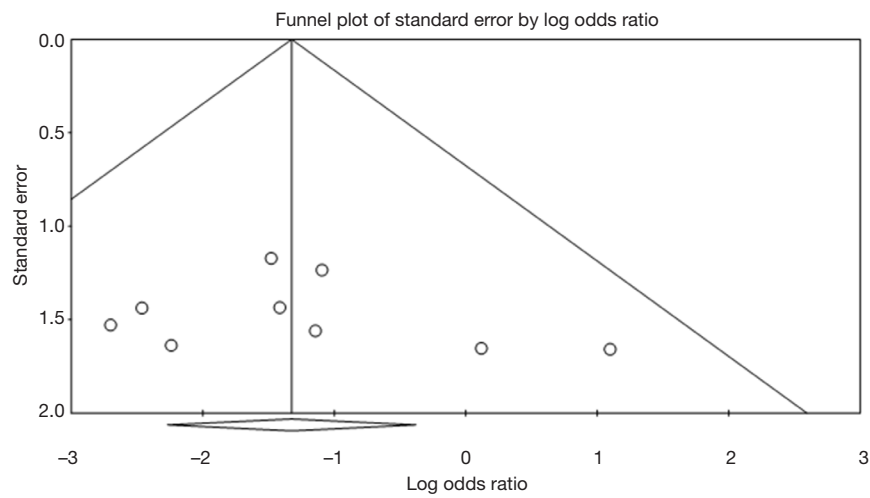


Figure S2 Funnel plot of studies reporting endocarditis rates in VSRR and CVG (Egger's test 1-tail $P=0.27$). VSRR, valve sparing root replacement; CVG, composite valve graft.

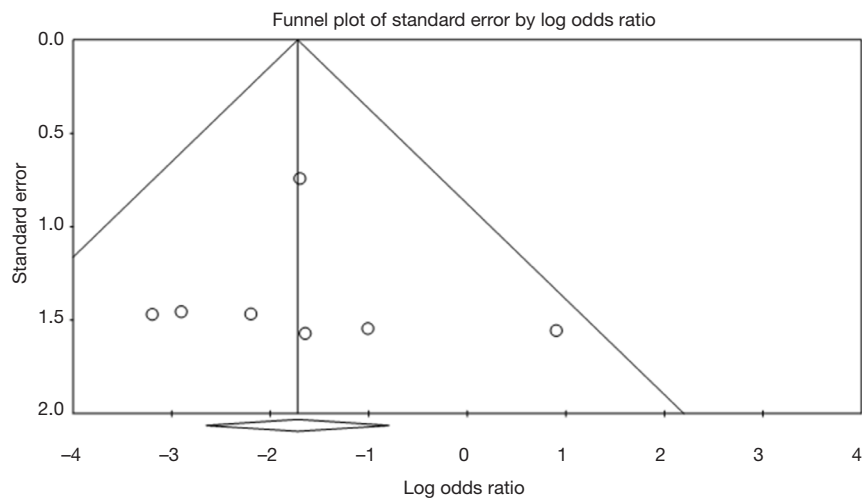


Figure S3 Funnel plot of studies reporting late bleeding events in VSRR and CVG (Egger's test 1-tail $P=0.45$). VSRR, valve sparing root replacement; CVG, composite valve graft.

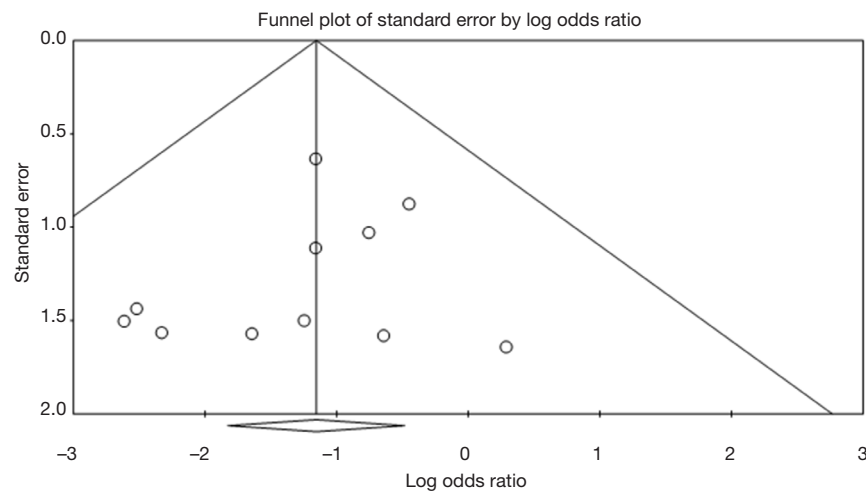


Figure S4 Funnel plot of studies reporting thromboembolic events in VSRR and CVG (Egger's test 1-tail $P=0.19$). VSRR, valve sparing root replacement; CVG, composite valve graft.

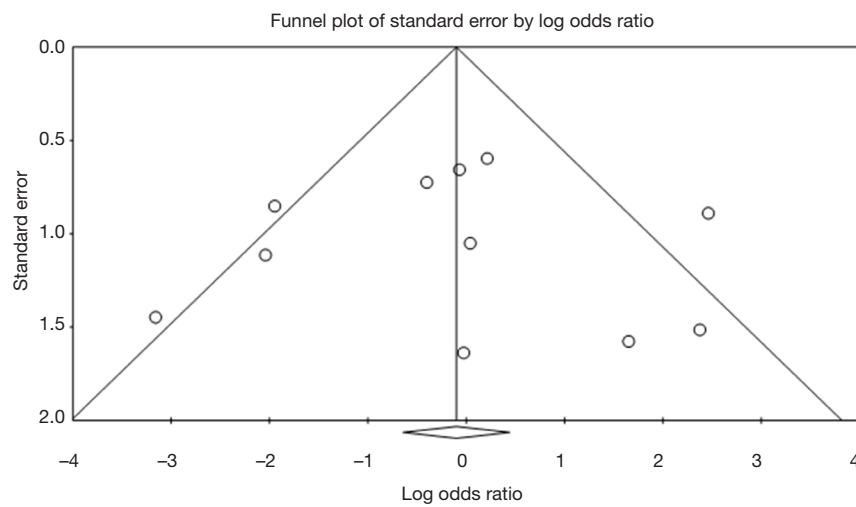


Figure S5 Funnel plot of studies reporting reintervention events in VSRR and CVG (Egger's test 1-tail $P=0.49$). VSRR, valve sparing root replacement; CVG, composite valve graft.

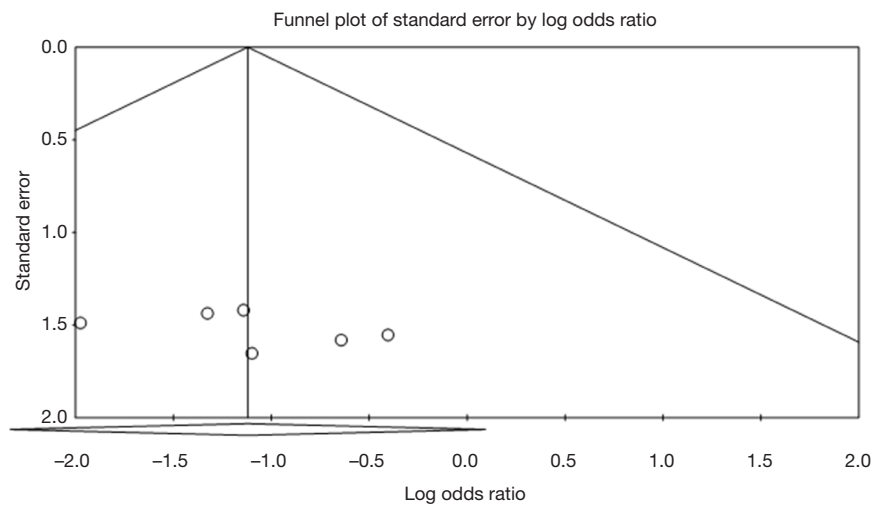


Figure S6 Funnel plot of studies reporting early mortality in VSRR and CVG (Egger's test 1-tail $P=0.21$). VSRR, valve sparing root replacement; CVG, composite valve graft.

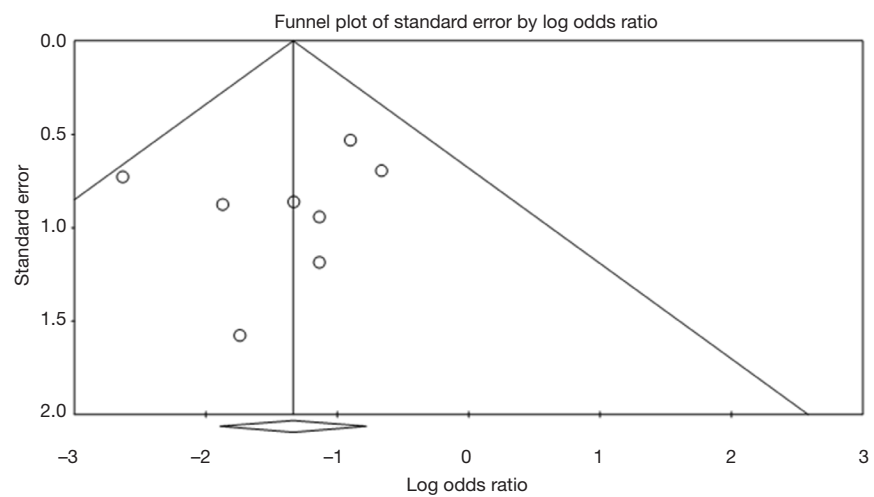


Figure S7 Funnel plot of studies reporting late mortality in VSRR and CVG (Egger's test 1-tail $P=0.27$). CVG, composite valve graft; VSRR, valve sparing root replacement.