Sildenafil for improving outcomes in patients with corrected valvular heart disease and persistent pulmonary hypertension: a multicenter, double-blind, randomized clinical trial

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Aims

We aimed to determine whether treatment with sildenafil improves outcomes of patients with persistent pulmonary hypertension (PH) after correction of valvular heart disease (VHD).

Methods and results

The sildenafil for improving outcomes after valvular correction (SIOVAC) study was a multicentric, randomized, parallel, and placebo-controlled trial that enrolled stable adults with mean pulmonary artery pressure ≥ 30 mmHg who had undergone a successful valve replacement or repair procedure at least 1 year before inclusion. We assigned 200 patients to receive sildenafil (40 mg three times daily, n = 104) or placebo (n = 96) for 6 months. The primary endpoint was the composite clinical score combining death, hospital admission for heart failure (HF), change in functional class, and patient global self-assessment. Only 27 patients receiving sildenafil improved their composite score, as compared with 44 patients receiving placebo; in contrast 33 patients in the sildenafil group worsened their composite score, as compared with 14 in the placebo group [odds ratio 0.39; 95% confidence interval (CI) 0.22–0.67; P < 0.001]. The Kaplan–Meier estimates for survival without admission due to HF were 0.76 and 0.86 in the sildenafil and placebo

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groups, respectively (hazard ratio 2.0, 95% CI = 1.0–4.0; log-rank P = 0.044). Changes in 6-min walk test distance, natriuretic peptides, and Doppler-derived systolic pulmonary pressure were similar in both groups.

**Conclusion**

Treatment with sildenafil in patients with persistent PH after successfully corrected VHD is associated to worse clinical outcomes than placebo. Off-label usage of sildenafil for treating this source of left heart disease PH should be avoided.

The trial is registered with ClinicalTrials.gov, number NCT00862043.

**Keywords**

Sildenafil • Pulmonary hypertension • Valvular heart disease • Heart failure

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**Introduction**

The most common cause of pulmonary hypertension (PH) worldwide is left heart disease (LHD), and valvular heart disease (VHD) is amongst the leading causes of this type of secondary PH. Pulmonary hypertension affects virtually all patients with severe symptomatic mitral valve disease and up to 65% of those with symptomatic aortic stenosis. Mitral and aortic valve diseases increase left atrial pressure which, in turn, leads to an initially passive and potentially reversible increase in pulmonary pressures. Vascular injury then triggers a cascade of venous and small artery remodelling, non-reversible arterial PH, and eventually, right ventricular dysfunction. Regression of PH is frequently incomplete after the correction of the valvular lesion, persisting in up to 75% of patients with moderate or severe preoperative PH. Furthermore, PH sometimes develops lately in patients who did not show PH before valve surgery. Once established, PH in corrected VHD is an untreatable risk-factor of mortality and disability in the long-term.

5-phosphodiesterase (PDE5) inhibitors have proven clinical efficacy in pulmonary arterial hypertension, but have shown discordant results in the field of LHD-PH. Nonetheless, sildenafil is frequently used off-label for treating this condition. In the setting of VHD, short-term studies have shown favourable effects of the drug in the immediate phases after surgery. To our knowledge no clinical trial has yet addressed the chronic effects of PDE5 inhibitors aimed specifically at treating persistent PH after correction of VHD. The sildenafil for improving outcomes after valvular correction (SIOVAC) trial was designed to test the hypothesis that, as compared with placebo, long-term therapy with the PDE5-inhibitor sildenafil improves clinical outcomes of patients with persistent PH after successful correction of the underlying VHD.

**Methods**

**Study design**

SIOVAC is an investigator-driven, academically sponsored, multicentric, randomized, double-blind, placebo-controlled, and parallel clinical trial. The study was performed in 18 academic hospitals in Spain, and the Fundación de Investigación Biomédica Hospital Gregorio Marañón served as the co-ordinating centre. The trial protocol (see Supplementary Material S2) was authorized by the Spanish Agency of Medicinal Products and Medical Devices and approved by the Reference Ethics Committee and the Local Ethics Committees of all participating institutions. All patients provided written informed consent. Randomization and clinical monitoring were performed by Chiltem International Ltd which also acted as the Data and Co-ordinating Centre in terms of study drug distribution and centralized data collection. An external adjudication and data safety monitoring board (ADSMB) reviewed all major adverse events and adjudicated clinical outcomes.

**Patients**

Patients were screened in outpatient clinics and imaging laboratories of the participating institutions (see Supplementary Material S1, Figure S1). Inclusion criteria for randomization were: (i) age older than 18 years, (ii) unequivocal demonstration of PH (a mean pulmonary arterial pressure ≥ 30 mmHg by catheterization within the 30 days prior to randomization), (iii) a successful surgical or percutaneous valvular replacement or repair procedure (leading to a complete correction of left heart valve disease and performed at least 1 year before inclusion), and (iv) a stable clinical condition (no changes in concomitant medication or hospital admissions for heart failure (HF) in the previous month). Major exclusion criteria were: (i) haemodynamically significant residual valvar or prosthetic valve dysfunction (patient-prosthesis mismatch or more than mild valvar or prosthetic valve stenosis or regurgitation, as assessed by the investigators according to current practice guidelines), (ii) systolic blood pressure < 90 mmHg, (iii) myocardial infarction, stroke, or life-threatening arrhythmia within the last 6 months, (iv) severe renal impairment (creatinine clearance <30 mL/min) or hepatic dysfunction, (v) life expectancy <2 years, or (vi) any established contraindication for sildenafil (see Supplementary Material S1, Table S1).

**Randomization and masking**

Patients were randomly assigned (1:1) to receive either sildenafil or placebo. Randomization was balanced using randomly permuted blocks of size four. Sites received the Investigational Product Kits containing two bottles of 550 tablets of the study drug with the patient’s treatment allocation code. Investigators and patients were masked to treatment assignment. Active treatment was re-bottled sildenafil (20 mg Revatio tablets, Pfizer), whereas the placebo manufacturing process ensured identical appearance to the active drug (see Supplementary Material S1).

**Procedures**

In patients in whom recent catheterization data was unavailable (88 patients, 44%) but showed a systolic pulmonary artery pressure ≥ 50 mmHg in a screening echocardiographic study, a per-protocol right heart catheterization procedure was performed. An acute vasoreactivity test with open-label sublingual sildenafil (100 mg) was performed for patients undergoing per-protocol catheterization. For patients who underwent catheterization prior to enrolment, vasoreactivity results were registered retrospectively whenever available. The diastolic and (mean) transmural pulmonary pressure gradients were calculated subtracting the mean...
pulmonary wedge pressure from the diastolic and mean pulmonary artery pressures, respectively.

Although the recommended dose of sildenafil for pulmonary arterial hypertension is 20 mg three times daily (t.i.d.), most clinical trials of sildenafil in LHD-PH have used higher doses.12–16,21,22 Accordingly, we chose a target study dose of 40 mg orally t.i.d. Patients with a low body-mass index, hypotension, or showing severe hypotension during the vasoactivity test initiated a 20 mg t.i.d. for 2 weeks.

Randomized patients underwent clinical assessment, 6-min walk test, and Doppler-echocardiography examinations at baseline, 3 and 6 months. Blood sampling for brain natriuretic peptide (BNP) measurements and magnetic resonance examinations (selected sites, in patients without contraindications) were performed at baseline and 6 months. All explorations were performed at least 4 h (half-life of sildenafil) after taking the study drug. Concomitant medication was recorded during clinical revisions and adherence was monitored by pill-counts at the 3 and 6 month visits. Blinded core laboratories analysed cardiac imaging and BNP concentrations.

**Outcomes**

The primary endpoint was based on the composite clinical score at 6 months. This score has demonstrated good sensitivity in clinical trials in the field of HF23 and fulfils the requirements for PH trials. The composite clinical score combines three elements: (i) major clinical events, defined as occurrence of death (of any cause) or hospital admission for HF requiring intravenous diuretic treatment with or without overnight stay, which is objective evidence of change in clinical status, (ii) World Health Organisation (WHO) functional classification, which relies on the physician assessment, and (iii) the patient global self-assessment, which relies on patient’s criteria. The self-assessment score is obtained interviewing the patient for his/her perception of change from his/her baseline clinical condition at enrollment.23 The composite clinical score classifies patient’s outcome in three categories: (i) worsened, if he/she presents a major clinical event, increases his/her WHO functional class, or self-reports a moderately or markedly worse category in the global-self-assessment, (ii) improved, if he/she has not suffered a major clinical event and his/her functional class has improved or reports moderate or marked improvement in global self-assessment or (iii) unchanged (otherwise). In case of discordant information, most objective events (death or HF admission) prevail over the change in functional class; the latter, in turn prevails over patient’s self-assessment. The ADSMB blindly adjudicated the composite clinical score in every patient.

Secondary endpoints were (i) the composite clinical score adjusted by co-variables (gender, age, and baseline WHO functional class), (ii) all-cause mortality, (iii) cardiovascular mortality, (iv) Kaplan–Meier analysis of major clinical events (as defined above), and (v) number of hospital admissions because of HF requiring intravenous diuretics. Other secondary endpoints were changes from entry to 6-month follow-up in (vi) WHO functional capacity, (vii) 6-min walk test distance, and (viii) plasma BNP levels. Imaging secondary endpoints were the change in systolic pulmonary pressure and in ventricular volumes at 6 months, by Doppler-echocardiography and magnetic resonance, respectively. Interactions between the primary endpoint and a number of baseline variables were pre-specified as exploratory analyses (see Supplementary Material S2).

**Statistical analysis**

The null hypothesis was that at the end of the 6-month follow-up period there is no difference between patients treated with placebo and sildenafil in the distribution of the three categories of the composite clinical score. The alternative hypothesis was that compared to placebo, sildenafil increases the proportion of patients who improve and decreases the proportion of patients who worsen their composite score. We used the mathematical formulation established for ordinal outcomes to calculate sample size.25 We initially estimated proportions of improved, worsened,
Table 1  Baseline characteristics

|                                | Sildenafil (n = 104) | Placebo (n = 96) | P-value |
|--------------------------------|----------------------|------------------|---------|
| Age (years), median (IQR)      | 70 (65, 77)          | 73 (67, 77)      | 0.23    |
| Women, n (%)                   | 76 (73)              | 78 (81)          | 0.23    |
| Weight (Kg), median (IQR)      | 66 (59, 78)          | 72 (62, 80)      | 0.13    |
| Body mass index (Kg m\(^{-2}\)), median (IQR) | 26.5 (24.0, 30.0)    | 28.4 (25.3, 32.4) | 0.04    |
| Systolic blood pressure (mmHg), median (IQR) | 131 (119, 144)       | 140 (127, 154)   | 0.02    |
| Diastolic pressure (mmHg), median (IQR) | 70 (64, 80)          | 70 (63, 80)      | 0.94    |
| Heart rate (beats min\(^{-1}\)), median (IQR) | 72 (67, 79)          | 70 (65, 82)      | 0.71    |
| Heart valve procedures         |                      |                  |         |
| Time from last valvular surgery (years), median (IQR) | 7.5 (4.2, 13.1)      | 5.8 (3.0, 12.3)  | 0.12    |
| Isolated mitral valve surgery, n (%) | 27 (26)              | 33 (34)          | 0.22    |
| Isolated aortic valve replacement, n (%) | 8 (8)                | 9 (9)            | 0.80    |
| Mitral and aortic valve surgery, n (%) | 29 (28)              | 16 (17)          | 0.06    |
| Mitral and tricuspid valve surgery, n (%) | 26 (25)              | 23 (24)          | 0.87    |
| Aortic and tricuspid valve surgery, n (%) | 0 (0)                | 1 (1)            | 0.48    |
| Mitral, aortic and tricuspid valve surgery, n (%) | 14 (14)              | 14 (15)          | 0.84    |
| Patients with re-interventions, n (%) | 39 (38)              | 24 (25)          | 0.07    |
| Coronary artery revascularization |                      |                  |         |
| Coronary artery bypass graft, n (%) | 3 (3)                | 10 (10)          | 0.06    |
| Percutaneous coronary intervention, n (%) | 5 (5)                | 7 (7)            | 0.66    |
| Cardiovascular risk factors    |                      |                  |         |
| Hypertension, n (%)            | 59 (57)              | 69 (72)          | 0.04    |
| Hyperlipidemia, n (%)          | 51 (49)              | 34 (35)          | 0.07    |
| Diabetes, n (%)                | 31 (30)              | 27 (28)          | 0.91    |
| Smoking, n (%)                 | 7 (7)                | 6 (6)            | 1.00    |
| Other comorbidities            |                      |                  |         |
| Atrial fibrillation, n (%)     | 77 (74)              | 77 (80)          | 0.39    |
| WHO functional classification  |                      |                  | 0.84    |
| I, n (%)                       | 8 (8)                | 8 (8)            |         |
| II, n (%)                      | 51 (51)              | 44 (46)          |         |
| III, n (%)                     | 42 (42)              | 43 (45)          |         |
| 6-min walk test distance (m), median (IQR) | 361 (285, 418)       | 342 (250, 382)   | 0.07    |
| Concomitant medications        |                      |                  |         |
| Acenocoumarol or warfarin, n (%) | 97 (93)              | 81 (84)          | 0.70    |
| Aspirin, n (%)                 | 11 (11)              | 11 (12)          | 1.00    |
| Diuretics, n (%)               | 89 (86)              | 84 (88)          | 0.99    |
| Aldosterone receptor antagonist, n (%) | 46 (44)              | 38 (40)          | 0.77    |
| Digoxin, n (%)                 | 43 (41)              | 41 (43)          | 1.00    |
| ACE inhibitors, n (%)          | 45 (43)              | 33 (34)          | 0.47    |
| Angiotensin II receptor blocker, n (%) | 22 (21)              | 20 (21)          | 1.00    |
| Beta-blocker, n (%)            | 53 (51)              | 43 (45)          | 0.69    |
| Calcium antagonist, n (%)      | 11 (11)              | 22 (23)          | 0.07    |
| Core laboratory biomarker data |                      |                  |         |
| BNP (pg mL\(^{-1}\)), median (IQR) | 63 (28, 166)         | 54 (25, 118)     | 0.40    |
| Cardiac catheterization data   |                      |                  |         |
| Right atrial pressure (mmHg), median (IQR) | 12 (9, 16)           | 12 (10, 17)      | 0.51    |
| Pulmonary artery oxygen saturation (%), median (IQR) | 64 (60, 70)         | 64 (57, 69)     | 0.38    |
| Mean pulmonary artery pressure (mmHg), median (IQR) | 39 (34, 46)         | 37 (34, 44)     | 0.25    |
| Mean wedge pulmonary pressure (mmHg), median (IQR) | 23 (19, 26)         | 22 (19, 26)     | 0.92    |
| Cardiac index (L s\(^{-1}\) m\(^{-2}\)), median (IQR) | 2.8 (2.4, 3.2)      | 2.8 (2.3, 3.4)  | 0.80    |
| Mean transpulmonary pressure gradient (mmHg), median (IQR) | 16.0 (13.0, 22.0)  | 15.0 (12.0, 20.0) | 0.35 |
| Diastolic transpulmonary pressure gradient (mmHg), median (IQR) | 2.0 (0.0, 6.0)      | 3.0 (0.0, 7.0)  | 0.44    |
| Pulmonary vascular resistance (Wood units), median (IQR) | 3.4 (2.4, 4.6)      | 3.1 (2.2, 4.9)  | 0.33    |

ACE, angiotensin-converting enzyme; BNP, brain natriuretic peptide; IQR, interquartile range; WHO, World Health Organisation.
and unchanged categories to be 15%, 20%, and 65%, respectively in the placebo group. We assumed an absolute 10% increase in the proportion of improvement in the sildenafil group \[\text{odds ratio (OR)} = 1.90\]. Using a two-sided level of significance of \(\alpha = 0.05\), these assumptions resulted in 322 patients needed for an 80% power to reject the null hypothesis. Estimating a 10% attrition rate, the initial sample size was 354 patients. Sample-size recalculation without unblinding was pre-specified after completing the follow-up of the first 100 patients. This analysis showed a higher than expected incidence of the worsened category in the global study population. After confirming no significant differences in the number of major clinical events between blinded groups, the ADSMB authorized to continue the study and requested to recalculate sample size. Using the observed proportions of the first 100 patients, the power to reject the null hypothesis was re-estimated in 190 analysable patients; based on the 4% attrition rate observed in the first 100 patients, the final sample size was re-adjusted to 198 patients.

The safety analysis set included all randomized patients who received at least one dose of the study drug. The full-analysis set (modified-intention-to-treat set) included all randomized patients who took at least one dose of the study drug, and on whom it was possible to evaluate the composite clinical score in at least one time-point. The per-protocol set excluded all patients with major protocol deviations.

We used an ordinal logistic regression model to calculate ORs for the primary endpoint under the proportionality assumption. Patients with an undetermined composite clinical score were excluded from the primary endpoint. However, sensitivity analyses were also pre-specified in which these patients’ outcomes were imputed as either ‘unchanged’ or using monotone logistic regression from baseline and 3 months variables. Odds ratios adjusted by age, sex, and WHO functional class were calculated as secondary endpoints. Time-to-event data was analysed using the Kaplan–Meier method, the log-rank test, and Cox regression. Quantitative secondary-endpoints were analysed using linear mixed-models for longitudinal data accounting for the fixed-effects of the visit, the treatment group and their interaction. Changes in functional class were analysed using a cumulative-link mixed-model for ordinal responses. Interaction analyses with baseline co-variables were performed using a logistic-regression model accounting for their interaction with the treatment group either continuously or by binary categorization. Signification was established as \(P\)-value \(<0.05\) (two-sided). Data analysis was performed by Chiltern International Ltd and the investigators using SAS software, version 9.2 (SAS Institute, Inc.) and R version 3.3.2. The study is registered with ClinicalTrials.gov NCT00862043 and EudraCT 2007-007033-40.

**Results**

From May 2009 to December 2015, 231 patients were enrolled, but 31 did not meet the mean pulmonary arterial pressure inclusion criterion (Figure 1). Thus, 200 patients were randomized to receive either sildenafil \((n = 104)\) or placebo \((n = 96)\). Three patients in the sildenafil and one in the placebo group abandoned the study without undergoing follow-up visits or reporting clinical events. Thus, the full analysis set consisted of 196 patients, 101 in the sildenafil, and 95 in the placebo group. The per-protocol set consisted of 162 patients, 80 receiving sildenafil, and 82 placebo. The study was completed by 170 patients \((85 \text{ in the placebo group and } 85 \text{ in the control group})\).

Nine patients took \(20 \text{ mg t.i.d.

Figure 2** Primary endpoint.** The composite clinical score accounts for the combination of death due to any cause, hospitalization due to heart failure requiring intravenous diuretic treatment, change in the World Health Organisation (WHO) functional class or relevant changes in the patient global self-assessment. Total bars show the proportion of patients in each category, stacked bars show the criterion used to adjudication to each category, and the table shows the number of patients. Data are shown for the full analysis set. Odds ratio calculated using ordinal logistic regression under the proportionality assumption.
persistent hypotension, and the other five upon investigators preferences.

Characteristics of the study patients

Most baseline clinical characteristics of randomized patients were not different between groups (Table 1). Median (IQR) age was 72 (66–77) years old, 154 were women, 154 were in atrial fibrillation, and 85 were in WHO functional Class III. Most patients (n = 182) had undergone mitral valve surgery (valve replacement in 160 and valve repair in 122), 91 had undergone aortic valve replacement, and 78 had undergone tricuspid valve surgery. Valvular interventions had been surgical in all patients except in 3 cases of transcatheter aortic valve replacement. Approximately one-third of patients had undergone repeated interventions. Baseline haemodynamic (Table 1) and imaging characteristics (see Supplementary Material S2, Table S2) were not significantly different between groups.
Primary endpoint

In the sildenafil group, only 27 patients improved their composite clinical score at 6 months as compared with 44 patients in the placebo group. By contrast, 33 patients in the sildenafil group worsened their primary outcome, as compared with 14 patients in the placebo group [OR for improvement 0.39; 95% confidence interval (CI) 0.22–0.67; \( P < 0.001 \)] (Figure 2). These unfavourable outcomes of patients taking sildenafil were confirmed in the per-protocol set (OR 0.42; 95% CI 0.24–0.76; \( P = 0.004 \)), as well as in the two sensitivity analyses in which the five patients with undetermined classifications were imputed, either as ‘unchanged’ (OR 0.40; 95% CI 0.23–0.68; \( P < 0.001 \)), or using monotonic logistic regression (OR 0.41; 95% CI 0.24–0.71; \( P = 0.001 \)).
Secondary endpoints

Unfavourable composite clinical scores in the sildenafil group were also confirmed when adjusting for co-variables such as age (OR 0.39; 95% CI 0.21–0.62; \( P < 0.001 \)), sex (OR 0.39; 95% CI 0.22–0.67; \( P < 0.001 \)), and baseline WHO functional class (OR 0.38; 95% CI 0.22–0.67; \( P < 0.001 \)). There were five deaths during the study, three in the sildenafil group (two of cardiac origin; one abdominal haemorrhage) and two in the placebo group (one cardiac; one pulmonary haemorrhage; log-rank \( P = 0.72 \)). The three cardiac deaths were due to HF (log-rank test \( P = 0.63 \) for sildenafil vs. placebo). The Kaplan–Meier estimates of survival at 6 months without major clinical events (death or hospitalization due to HF) were 0.76 (95% CI 0.68–0.85) and 0.86 (95% CI 0.78–0.94), respectively (hazard ratio 2.0, 95% CI 1.0–4.0; log-rank test \( P = 0.04 \)). The Kaplan–Meier estimates of survival at 6 months without major clinical events (death or hospitalization due to HF) were 0.76 (95% CI 0.68–0.85) and 0.86 (95% CI 0.78–0.94), respectively (hazard ratio 2.0, 95% CI 1.0–4.0; log-rank test \( P = 0.04 \)). There were 31 HF hospital admissions requiring intravenous diuretics in the sildenafil group vs. 22 in the placebo group (OR 0.43, 95% CI 0.20–0.94; \( P = 0.035 \); Figure 3A). There were 31 HF hospital admissions requiring intravenous diuretics in the sildenafil group vs. 22 in the placebo group (OR 0.43, 95% CI 0.20–0.94; \( P = 0.035 \); Figure 3A). There were no significant differences between groups in the changes from baseline to 6 months in functional capacity, 6-min walk distance, BNP levels, or systolic pulmonary artery pressure (Figure 3C–F; see Supplementary Material S1, Tables S3 and S4). Magnetic resonance data showed LV dilatation from baseline to month 6 only in the sildenafil group, resulting in significant differences in the changes of LV end-diastolic and end-systolic volumes between groups (\( P = 0.04 \) and 0.05, respectively; see Supplementary material online, Figure S1).

Pre-specified subgroup analyses

Binary interaction analyses did not identify any pre-specified baseline co-variable suggesting a benefit for the active treatment (Figure 4). Furthermore, no predictive value for response was identified for acute vasoreactivity data. Quantitative interactions analyses also failed to suggest a potential benefit of treatment in any range of the tested haemodynamic variables (Figure 5).

Adverse events

The sildenafil group showed a non-significant trend towards more frequent investigator-reported serious adverse events than the placebo group, in particular related to the study drug (Table 2). More frequent infectious adverse events were observed in the placebo group, in particular related to the study drug (Table 2). More frequent infectious adverse events were observed in the placebo group, in particular related to the study drug (Table 2). More frequent infectious adverse events were observed in the placebo group, in particular related to the study drug (Table 2).

Discussion

To the best of our knowledge this is the first clinical trial targeted to persistent PH in patients with corrected VHD. Contrary to our alternative hypothesis, long-term treatment with oral sildenafil negatively impacted outcome compared with placebo. These data confirm the recommendation of current practice guidelines against using PDE5 inhibitors and other drugs approved for pulmonary arterial hypertension in patients with LHD-PH.
Because it is believed to be safe and well tolerated, sildenafil is frequently used off-label to treat LHD-PH. The striking acute effects of the drug described in patients with native aortic valve stenosis, and after left-side valvular surgery have probably further expanded its use in VHD. Noticeably, our vasoreactivity test also lowered pulmonary pressures and increased cardiac output (Supplementary material online, Table S1, Table 1), in identical direction to these acute studies. Thus, the favourable acute haemodynamic effects of sildenafil may not be predictive of long-term outcome after prolonged administration in patients with LHD-PH.

Previous evidence of chronic treatment with PDE5 inhibitors in LHD-PH is controversial. Small clinical trials have suggested that sildenafil may improve the haemodynamic profile, overall exercise performance, and quality of life of patients with chronic LHD-PH of non-valvular etiology. Favourable evidence is highest in patients with HF and reduced ejection fraction, but also positive results have been reported in selected patients with HF and preserved ejection fraction. More recent clinical trials have questioned the clinical efficacy of sildenafil in patients with HF, and neutral findings have been reproduced with riociguat, another drug targeted to the nitric oxide signalling pathway.

Most clinical trials of sildenafil in the field of LHD-PH have excluded elderly patients, and women have been frequently underrepresented or formally excluded. In fact, sildenafil resulted in a higher pulmonary capillary pressure than placebo in the only HF clinical trial showing demographic and comorbidity patterns comparable to our study. In our trial, neutral changes in 6-min walk test distance and natriuretic peptides did not match the incidence of hard events (readmission for HF), underscoring the need of using clinical outcomes in future trials in the field.

Although the mechanisms leading to worse outcomes of patients taking sildenafil in our study are necessarily speculative, a chronic increase in pulmonary capillary pressure is the most plausible explanation. The combination of advanced age, prevalent atrial fibrillation, and long-standing atrial overload, reduces atrial compliance in patients with VHD. In this context, pulmonary arterial vasodilation may rise capillary pressure because the left heart is not able to accommodate the increase in right ventricular output. This would render capillary pressure particularly sensitive to an increase in flow, predisposing to HF decompensation. Remarkably, our imaging findings support a certain degree of volume overload induced by sildenafil. Further analyses are need to clarify why drug-induced ventricular dilatation was higher for the LV than the RV.

As in other trials with sildenafil in HF, we did not exclude patients based on specific haemodynamic patterns of PH. It can be speculated that according to current guidelines, no benefit of...
sildenafil would to be expected if the SIOVAC population was mostly comprised by patients with isolated post-capillary PH. However, 57% of patients included in our trial showed a pulmonary vascular resistance > 3 Wood units, compatible with combined post- and pre-capillary PH. Although limited due to sample size, interaction analyses showed no evidence of potential benefit in any specific baseline or vasoreactivity haemodynamic profile. Thus, we believe our study adds clarifying information on the negative role of pulmonary vasodilators in LHD-PH.

Equivalent mortality rate in our study was 5% per year. This figure is similar to the expected mortality of patients with pulmonary arterial hypertension who share the functional, biomarker, and haemodynamic profiles of patients in our study. Thus, VHD-PH should not be conceived as a benign condition and further basic and clinical research should continue to explore alternative therapies in this field.

**Study limitations**

The study group was heterogeneous in terms of VHD primary lesions, but sample size did not allow for a more detailed subgroup comparison of treated valves or type of surgery for example repair or replacement. The study was also underpowered to obtain significant results in most of the secondary endpoints. The composite score on which we based our primary endpoint merges outcomes of diverse significance and may seem subjective. However, composite scores meet current consensus of incorporating multiple outcome measures, circumvent the need of an allocation for testing multiple hypothesis, avoid the problems of competing risks and, most importantly, allow for lowering the cost of clinical trials by reducing required sample sizes. Furthermore, composite clinical scores which include self-assessment scales have been useful to demonstrate the efficacy of pharmacological and non-pharmacological therapies in HF. In addition, disaggregated analyses demonstrated that differences in the primary endpoint were due to an increase of the risk of HF decompensation in patients taking sildenafil. Thus, we believe the study provides reliable evidence on the impact of sildenafil on clinical outcomes of patients with LHD-PH due to VHD.

**Conclusions**

Treatment with oral sildenafil 40 mg t.i.d. for 6 months in patients with persistent PH after successful correction of VHD is associated to unfavourable clinical outcomes as compared to placebo. In the light this study, open label use of sildenafil in PH due to VHD should be discouraged, in agreement with current PH practice guidelines. Further efforts to identify novel therapeutic targets in this particular source of PH are needed.

**Supplementary material**

Supplementary material is available at European Heart Journal online.

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**Conflict of interest**: none declared.

**References**

The list of references is available on the online version of this paper.