# National Institute for Health and Care Excellence

**Draft** 

# Heart valve disease presenting in adults: investigation and management

[C] Evidence reviews for pharmacological management

NICE guideline

Intervention evidence review underpinning recommendations 1.2.1 and 1.2.2 and research recommendations in the NICE guideline

March 2021

**Draft for Consultation** 

This evidence review was developed by the National Guideline Centre



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### 1<sub>1</sub> Introduction

- 2 The management of heart valve disease necessitates mechanical intervention. However,
- 3 medical management may also play a role in the management of heart failure symptoms,
- 4 particularly in heart failure with consequent secondary heart valve disease but also in primary
- 5 heart valve disease with consequent valvular heart failure awaiting valve intervention. In the
- 6 absence of heart failure, heart valve disease may impact the medical management of
- 7 coexistent conditions, for example systemic hypertension. Furthermore, attempts to treat or
- 8 slow down the progression of heart valve disease were made over the years with a variety of
- 9 drugs. Consequently, it is important to determine the clinical and cost effectiveness of
- 10 medical management in adults with heart valve disease with and without concomitant heart
- 11 failure.

## 2 Pharmacological management

2.13 Review question: In adults with heart valve disease without

- 4 concomitant heart failure, what is the clinical and cost
- 5 effectiveness of alpha-blockers, angiotensin-converting
- 6 enzyme (ACE) inhibitors, angiotensin-II receptor blockers
- 7 (ARBs), beta blockers, calcium channel blockers, digoxin,
- 8 diuretics, statins and nitrates to improve clinical outcome?

#### 2.1.9 PICO table

2

10 For full details see the review protocol in Appendix A.

#### 11 Table 1: PICO characteristics of review question

able 1: PICO cr	naracteristics of review question
Population	Adults aged 18 years and over with diagnosed heart valve disease of at least moderate severity stratified by type:  Primary aortic [including bicuspid] stenosis  Primary aortic regurgitation  Primary mitral stenosis  Primary mitral regurgitation  Primary tricuspid regurgitation  Secondary heart valve disease – mitral regurgitation or tricuspid regurgitation  A study will be considered to cover a population with heart valve disease without concomitant heart failure if it meets all of the following criteria:  Diagnosis of native heart valve disease  Asymptomatic or have only very mild/low-level symptoms that would not affect daily life (this would include those reported to be in class I of the NYHA classification)  A normal LVEF
Interventions	<ul> <li>Alpha blockers</li> <li>Angiotensin-converting enzyme (ACE) inhibitors</li> <li>Angiotensin-II receptor blockers (ARBs)</li> <li>Beta blockers</li> <li>Calcium channel blockers</li> <li>Digoxin</li> <li>Diuretics</li> <li>Statins</li> <li>Nitrates (including nitroprusside)</li> <li>Any combination of 2 or more of the above</li> </ul>
Comparison(s)	Placebo or no treatment (usual care)     Other petitive comparator listed chave including combinations.
	<ul> <li>Other active comparator listed above, including combinations</li> </ul>

- All-cause mortality at ≥12 months (dichotomous)
- Cardiac mortality at ≥12 months (dichotomous)
- Health-related quality of life at 6 months and ≥12 months (continuous)
- Onset of symptoms or progression in NYHA class at ≥12 months
- Evidence of HVD progression on imaging (worsening of disease severity) at ≥ 12 months (dichotomous)
- Need for heart valve intervention (surgical or transcatheter) at ≥12 months (dichotomous)

Secondary outcomes (important outcomes):

- Exercise tolerance reported as any of the following (in order of relevance) at 12 months:
  - Supine bicycle workload (watts or % difference from predicted watts)
  - Treadmill exercise time (duration)
  - o Oxygen consumption on exercise testing (VO<sub>2</sub> max)
  - o Time to near maximal dyspnoea
  - o 6-minute walk test
  - Borg dyspnoea index

(Continuous)

 Withdrawal from the trial due to adverse events at 6 and 12 months (dichotomous)

Study design

RCTs or systematic reviews of RCTs

#### 2.1.2 Clinical evidence

#### 2.1.22 Included studies

4

- 3 Seventeen studies from twenty-seven papers were included in the review;<sup>3, 13, 14, 20, 21, 25-28, 36,</sup>
  - 41, 49, 58-60, 66, 71, 74, 104, 129, 131, 132, 143, 145, 146, 154, 177 these are summarised in Table 3 below.
- 5 Evidence from these studies is summarised in the clinical evidence summaries below (Table
- 4 to Table 15). Studies identified investigated pharmacological management in people with
- 7 aortic stenosis, aortic regurgitation and mitral regurgitation.
- 8 The identified studies included the following comparisons for each population stratum, with
- 9 some studies reporting more than one comparison:
- 10 Aortic stenosis:
- ACE inhibitors compared to placebo: 1 study<sup>21</sup>
- Beta blockers compared to placebo: 1 study<sup>66</sup>
- Diuretics compared to placebo: 1 study<sup>154</sup>
- Statins compared to placebo: 4 studies (14 papers)<sup>14, 25-28, 36, 41, 58-60, 71, 74, 131, 132</sup>
- 15 Primary aortic regurgitation
- ACE inhibitors compared to placebo/no treatment: 2 studies<sup>49, 177</sup>
- ACE inhibitors compared to calcium channel blockers: 2 studies<sup>13, 49</sup>
- ARBs compared to beta blockers: 1 study<sup>129</sup>
- Beta blockers compared to placebo: 1 study<sup>20</sup>
- Calcium channel blockers compared to placebo/no treatment: 2 studies<sup>49, 145</sup>

- Digoxin compared to calcium channel blockers: 1 study<sup>145</sup>
- 2 Primary mitral regurgitation
- ACE inhibitors compared to placebo: 3 studies<sup>104, 143, 177</sup>
- Beta blockers compared to placebo: 1 study<sup>4</sup>
- 6 No relevant RCTs investigating pharmacological management in people without concomitant
- 7 heart failure and mitral stenosis or tricuspid regurgitation were identified. No relevant studies
- 8 investigating the use of alpha blockers or nitrates were identified. No relevant studies
- 9 investigating the use of combinations of treatment were identified.
- 10 See also the study selection flow chart in Appendix C:, study evidence tables in Appendix D:,
- 11 forest plots in Appendix E:and GRADE tables in Appendix F:.

#### 2.1.22 Excluded studies

5

13 See the excluded studies list in Appendix I:.

1

# 2.1.2.3 Summary of clinical studies included in the evidence review

Table 3: Summary of studies included in the evidence review

Study	Intervention and comparison	Population	Outcomes	Comments
Ahmed 2012 <sup>4</sup>	Beta blockers (n=19) Oral metoprolol (Toprol XL) for 2 years. Starting dose of 12.5-25mg/day titrated up to the maximum tolerable dose at 2-week intervals. Maximum dose: 100mg/day.  Placebo (n=19) Oral placebo  No information on concurrent medication/care.	Primary mitral regurgitation (N=38) Severity of disease: Not stated Mechanism of disease: Degenerative  Defined by echocardiography (colour flow Doppler imaging).  Age (mean [SD]): 52.9 (9.1) years Presence or absence of uncontrolled systemic hypertension (140/85mmHg) at the end of the trial: Not stated/unclear	All-cause mortality at 2 years Cardiac mortality at 2 years Need for heart valve intervention at 2 years Serious adverse events at 2 years	Equipment/drugs provided by industry (AstraZeneca).  Study aim: To complete MRI analysis of the effects of treatment on left ventricular remodelling and function in people with chronic, isolated mitral regurgitation
Banaszewski 1998 <sup>13</sup>	Calcium-channel blockers (CCB) (n=12) Oral nifedipine 10-20mg three times a day for 2.75 years. Mean daily dose of 40mg.  Angiotensin-converting enzyme (ACE) inhibitors (n=13) Oral captopril 12.5-30mg three times a day for 2.75 years. Mean daily dose of 75mg.	Primary aortic regurgitation (N=31) Severity of disease: Moderate to severe (AR grade range: 2-4) Mechanism of disease: Not stated  Defined by echocardiography and cardiac catheterisation.	Onset of symptoms or progression of NYHA class at 2.75 years Evidence of HVD progression on imaging (worsening of disease severity) at 2.75 years	The study included an acute phase where participants underwent cardiac catheterisat and exercise therapy after a single dose of nifedipine, which was then repeated after 24 hours, a long term (randomised phase of the study was started Study aim: To look at the short term haemodynamic effects of

Study	Intervention and comparison	Population	Outcomes	Comments
	Concurrent medication/care: The study only included people who were not using ACE inhibitors, calcium-channel blockers, diuretics, beta blockers and digitalis prior to the study.	Age (mean [SD]): 34.9 (10.1) years Presence or absence of uncontrolled systemic hypertension (140/85mmHg) at the end of the trial: Not stated/unclear (initial blood pressure was not hypertensive).		nifedipine and captopril on left ventricular volume responses, both at rest and at peak exercise, in asymptomatic people with moderate to severe isolated, untreated aortic regurgitation, enlarged left ventricle and normal left ventricular function, as well as long term effects of the two drugs in this population.
Broch 2016 <sup>20</sup>	Beta blockers (n=37) Oral metoprolol CR/XL 25mg doubled every week up to a target daily dose of 200mg or the maximum tolerable dose. Maintained for 6 months.  Placebo (n=38) Oral placebo  Concurrent medication/care: All people were allowed to use other vasoactive drugs (6 in each group using ACE inhibitors/ARBs, 2 in beta blocker arm using calcium- channel blockers, 3 in placebo arm using calcium-channel blockers, 5 in both arms using statins, 8 in beta blocker arm using acetylsalicylic acid, 2 in placebo arm using acetylsalicylic acid).	Primary aortic regurgitation (N=75) Severity of disease: Severe (Vena contracta width 7.6 (1.6)cm) Mechanism of disease: Not stated  Defined by cardiac magnetic resonance imaging and echocardiography. 55 (73%) of people had bicuspid aortic valves.  Age (mean [SD]): 44 (14) years Presence or absence of uncontrolled systemic hypertension (140/85mmHg) at the end of the trial: Absent (final blood pressure <140/85mmHg)	Quality of life at 6 months Exercise tolerance at 6 months	Equipment/drugs provided by industry (AstraZeneca)  Aim of the study: To examine the effect of metoprolol in asymptomatic people with chronic, moderate-to-severe aortic regurgitation, hypothesising the beta-blockade would reverse LV remodelling in these patients.

Study	Intervention and comparison	Population	Outcomes	Comments
Bull 2015 <sup>21</sup>	Angiotensin-converting enzyme (ACE) inhibitors (n=50) Oral ramipril 2.5mg daily for 2 weeks, raised to 5mg daily until the 3 month follow up, raised to 10mg daily for the rest of the study or until maximal tolerable dose (?rationale for larger dose).  Placebo (n=50) Oral placebo  Concurrent medication/care: Not stated	Primary aortic [including bicuspid] stenosis (N=100) Severity: Moderate or severe Mechanism of disease: Not stated  Defined by cardiac magnetic resonance imaging and echocardiography.  Age (mean [SD]): 68.57 (14.22) years Disease mechanism for aortic and mitral stenosis: Not stated/unclear Presence or absence of uncontrolled systemic hypertension (140/85mmHg) at the end of the trial: Not stated/unclear (at the start of the study 11 people in the ACE inhibitor arm and 18 people in the placebo arm had hypertension).	Need for heart valve intervention at 12 months Exercise tolerance at 12 months Withdrawal due to adverse events at 12 months	RIAS study Academic or government funding (Heart Research UK, and the Oxford Comprehensive Biomedical Research Centre, funded by the National Institute of Health Research).  Aims of the study: 1) To examine changes in myocardial physiology, in particular the regression of left ventricular mass, as well as other left ventricular physiological parameters using multiparametric cardiac magnetic resonance in people with moderate to severe aortic stenosis. 2) to assess the safety and tolerability of ramipril in these people. 3) to examine potential improvements in effort tolerance.
Chan 2010 <sup>27</sup> Subsidiary papers: Chan 2011 <sup>25</sup> Chan 2010 <sup>26</sup> Chan 2007 <sup>28</sup>	Statins (n=136) Oral rosuvastatin 40mg once a day for 3.5 years.  Placebo (n=136) Oral placebo  Concurrent medication/care: Not stated	Primary aortic [including bicuspid] stenosis (N=272) Severity of disease: Mild-to-moderate (mean peak aortic velocity was 3.16 (0.42) in the intervention arm and 3.19 (0.42) in the control arm, considered moderate severity in the British Society of Echocardiography guidelines)	All-cause mortality at 3.5 years Cardiac mortality at 3.5 years Need for heart valve intervention at 3.5 years Withdrawal due to adverse events at 3.5 years	ASTRONOMER study Study funded by industry (AstraZeneca Canada)  Aim of the study: To assess the effect of intensive lipid lowering with rosuvastatin on the progression of AS in asymptomatic people with mild to moderate AS and to assess the impact of intensive lipid lowering

Study	Intervention and comparison	Population	Outcomes	Comments
		Defined by echocardiography.  Age (mean [SD]): 57.9 (13.6) years Disease mechanism for aortic stenosis: Mixed (50% of people had bicuspid aortic valve disease, otherwise not stated).  Presence or absence of uncontrolled systemic hypertension (140/85mmHg) at the end of the trial: Not stated/unclear (initial blood pressure was not hypertensive).  Presence of coronary artery disease: No (people with coronary artery disease or any other indication for statins [apart from presence of hypercholesterolaemia] were excluded)		on adverse outcomes related to AS.
Cowell 2005 <sup>36</sup> Subsidiary papers: Houslay 2006 <sup>74</sup>	Statins (n=77) Oral atorvastatin 80mg once a day for 25 months. Concurrent medication/care: 43 taking aspirin, 12 taking ACE inhibitors, 21 taking beta blockers, 8 taking warfarin.  Placebo (n=78) Oral placebo	Primary aortic [including bicuspid] stenosis (N=155) Severity of disease: Severe (aortic jet velocity of at least 2.5m/s)  Defined by echocardiography.	Cardiac mortality at 25 months Onset of symptoms or progression of NYHA class at 25 months Need for heart valve intervention at 25 months Withdrawal due to adverse events at 25 months	SALTIRE study Funding not stated  Aim of the study: To establish whether intensive lipid-lowering therapy with atorvastatin would halt the progression or induce regression of aortic jet velocity on Doppler echocardiography, and of the aortic-valve calcium score on

Study	Intervention and comparison	Population	Outcomes	Comments
	Concurrent medication/care: 40 taking aspirin, 14 taking ACE inhibitors, 27 taking beta blockers, 12 taking warfarin.	Age (mean [SD]): 68 (10.5) years Disease mechanism for aortic stenosis: Calcific Presence or absence of uncontrolled systemic hypertension (140/85mmHg) at the end of the trial: Not stated/unclear (initial values were >140mmHg systolic but <85mmHg diastolic). Presence of coronary artery disease: 39 people had coronary artery disease. 20 had cerebrovascular disease.		computed tomography (CT), in people with calcific aortic stenosis.
Dichtl 2008 <sup>41</sup>	Statins (n=25) Oral atorvastatin 20mg once a day for 3-5 years. Concurrent medication/care: 7 people taking aspirin, 6 people taking ACE inhibitors, 1 person taking a calcium-channel blocker, 1 person taking a beta blocker, 1 person taking a vitamin K antagonist.  Placebo (n=25) Oral placebo  Concurrent medication/care: 14 people taking aspirin, 11 people taking ACE inhibitors, 2 people taking calcium-channel blockers, 5 taking beta	Primary aortic [including bicuspid] stenosis (N=50) Severity of disease: ?Severe (mean systolic gradients of ≥15mmHg and valvular stenosis flow velocities of ≥2.0m/s.  Defined by transthoracic echocardiography.  Age (mean [SD]): 67.0 (11.7) years Disease mechanism for aortic stenosis: Calcific Presence or absence of uncontrolled systemic hypertension (140/85mmHg) at the end of the trial: Not	All-cause mortality at 5 years Cardiac mortality at 5 years Need for heart valve intervention at 5 years Withdrawal due to adverse events at <6 months	Equipment/drugs provided by industry (Pfizer Austria)  Aim of the study: To further evaluate risk factors, the progression rate of disease, and possible beneficial effects of newonset lipid lowering therapy with atorvastatin at a standard daily dose of 20mg compared to placebo

Study	Intervention and comparison	Population	Outcomes	Comments
	blockers, 3 taking vitamin K antagonists.	stated/unclear (states 9 people in the intervention arm and 14 in the control arm had hypertension. Antihypertensive medication as prescribed for these people). Presence of coronary artery disease: 13 people had coronary artery disease (5 more in the atorvastatin arm than the control arm)		
Evangelista 2005 <sup>49</sup>	Calcium-channel blockers (CCB) (n=32) Oral nifedipine 20mg every 12 hours for 7 years  Angiotensin-converting enzyme (ACE) inhibitors (n=32) Oral enalapril 20mg daily for 7 years  No treatment (n=31)  Concurrent medication/care not stated	Primary aortic regurgitation (N=95)  Severity of disease: Severe Mechanism of disease: Not stated  Defined by physical examination, echocardiography, 12-lead electrocardiogram, chest radiography and radionuclide angiography at rest.  40 people had bicuspid aortic valve disease.  Age (mean [SD]): 44.35 (13.19) years Presence or absence of uncontrolled systemic hypertension (140/85mmHg) at the end of the trial:	All-cause mortality at 7 years Cardiac mortality at 7 years Onset of symptoms or progression of NYHA class at 7 years Evidence of HVD progression on imaging (worsening of disease severity) at 7 years Need for heart valve intervention at 7 years Withdrawal due to adverse events at 7 years	Academic or government funding (supported by a grant from the Red de Investigación Cooperativa de las Enfermedades Cardiovasculares from the Instituto de Salud Carlos III, Ministerio de Sanidad y Consumo, Spain)  Aim of the study: To ascertain whether either nifedipine or enalapril reduces or delays the need for valve surgery and whether these drugs exert any effect on the size and function of the left ventricle in this population

Study	Intervention and comparison	Population	Outcomes	Comments
		Present (final systolic blood pressure >140mmHg, final diastolic blood pressure mixed)		
Hansson 2017 <sup>66</sup>	Beta blockers (n=20) Oral extended-release metoprolol from 50mg up to a target daily dose of 200mg (over a six week titration period) or maximal dose without symptoms. Maintained for 5 months.  Placebo (n=20) Oral placebo  Concurrent medication/care: Not stated	Primary aortic [including bicuspid] stenosis (N=40) Severity of disease: Moderate-to-severe  Defined by echocardiography and cardiac magnetic resonance imaging.  Age (mean [SD]): 70.0 (5.1) years Disease mechanism for aortic stenosis: Not stated/unclear (7 people had bicuspid aortic valves, otherwise not stated). Presence or absence of uncontrolled systemic hypertension (140/85mmHg) at the end of the trial: Mixed (mean blood pressure metoprolol: 136/79 (13/8), mean blood pressure placebo: 140/81 (12/7)).	Quality of life at 5 months Exercise tolerance at 5 months Withdrawal due to adverse events at 5 months	Academic or government funding (Funded by the Lundbeck foundation, the Arvid Nilssons Foundation, the Health Research Fund of Central Denmark Region, Karen Elise Jensens Foundation, and Snedkermester Sophus Jacobsen and Hustru Astrid Jacobsens Foundation).  Aim of the study: To investigate whether metoprolol could improve myocardial efficiency (investigating the safety, haemodynamic and metabolic effects of metoprolol)
Marcotte 1997 <sup>104</sup>	Angiotensin-converting enzyme (ACE) inhibitors (n=12) Oral lisinopril 5mg for two weeks, then doubled every two weeks until maximal dose of	Primary mitral regurgitation (N=23) Severity of disease: At least moderate Mechanism of disease: Organic	All-cause mortality at 1 year Cardiac mortality at 1 year Quality of life at 6 months Quality of life at 1 year Exercise tolerance at 1 year	Study funded by industry (Merck Frosst Canada inc.)  Population may include people with congenital mitral regurgitation.

Study	Intervention and comparison	Population	Outcomes	Comments
	20mg a day or maximal tolerable dose. Maintained for 1 year.  Placebo (n=11) Oral placebo  Concurrent medication/care: No other cardiovascular medications	Defined by echocardiography.  Age (mean [SE]): 53.3 (2.4) years  Presence or absence of uncontrolled systemic hypertension (140/85mmHg) at the end of the trial: Not stated/unclear (absent at the start of the study).	Withdrawal due to adverse events at 12 months	Aim of the study: To determine the effectiveness of lisinopril in reducing the severity of mitral regurgitation in the population and ultimately in altering favourably the natural history of the disease
Roberts 2018 <sup>129</sup>	Angiotensin-II receptor antagonists (ARBs) (n=17) Oral losartan up-titrated to a maximum of 100mg per day for 1-3 weeks.  Beta blockers (n=17) Oral metoprolol CR to a maximum dose of 190mg for 1-3 weeks.  Concurrent medication/care: People were allowed to use normal antihypertensive medicines, which were then down-titrated or withdrawn completely while taking the drug.	Primary aortic regurgitation (N=46) Severity of disease: Severe (regurgitant volume 57.6 [35.8]mL) Mechanism of disease: Not stated  Defined by echocardiography.  Age (mean [SD]): 51.0 (14.1) years Presence or absence of uncontrolled systemic hypertension (140/85mmHg) at the end of the trial: Absent (blood pressure ranged between 117-118/63-69mmHg).	Exercise tolerance at 3 weeks	Academic or government funding (funded by the Health Research Council of New Zealand)  Cross-over study (0 day washout period).  Aim of the study: To compare the effects of losartan and metoprolol on aortic regurgitant fraction, LV and aortic function at rest and during exercise in asymptomatic people with chronic aortic regurgitation using cardiac magnetic resonance imaging.

Heart valve disease: DRAFT Pharmacological management

FOR CONSULTATION

Study	Intervention and comparison	Population	Outcomes	Comments
		Presence of coronary artery disease: No (people with coronary artery disease, cerebrovascular disease, peripheral arterial disease and diabetes mellitus were excluded).		
Sampaio 2005 <sup>143</sup>	Angiotensin-converting enzyme (ACE) inhibitors (n=27) Oral enalapril 5mg twice a day, titrated up to the maximal tolerated dose of at most 20mg twice a day (increased to 10mg at 2 weeks and 20mg at 4 weeks). Maintained for 1 year.  Placebo (n=27) Oral placebo  Concurrent medication/care: Not receiving therapy with any other vasodilators.	Primary mitral regurgitation (N=47) Severity of disease: Moderate to severe Mechanism of disease: Secondary to mitral prolapse or rheumatic heart disease  Defined by echocardiography.  Age (mean [SD]): 39 (15) years Presence or absence of uncontrolled systemic hypertension (140/85mmHg) at the end of the trial: Absent (final blood pressure ACE inhibitors: 122/78 (12/9)mmHg; final blood pressure placebo: 126/79 (12/8)mmHg).	Onset of symptoms or progression of NYHA class at 12 months Need for heart valve intervention at 12 months Exercise tolerance at 12 months	Academic or government funding (E.J. Zerbini foundation, São Paulo, Brazil)  At the start of the study 20 people were NYHA class I, 17 people were NYHA class II.  Aim of study: To evaluate the effects of enalapril on LV dimensions, LV systolic index, and functional capacity, with cardiopulmonary testing, after 12 months of therapy.
Scognamiglio 1990 <sup>145</sup>	Calcium-channel blockers (CCB) (n=38) Oral nifedipine 20mg twice	Primary aortic regurgitation (N=72) Severity of disease: Severe	Need for heart valve intervention at 1 year Withdrawal due to adverse	Funding not stated  Aim of the study: To verify
	daily for 1 year.		events at 1 year	whether long-term vasodilator therapy with nifedipine reduces

Study	Intervention and comparison	Population	Outcomes	Comments
	Placebo (n=34) Oral placebo  Concurrent medication/care: No cardioactive therapies.	Mechanism of disease: Not stated  Defined by Doppler colour flow imaging and confirmation by cardiac catheterisation.  Age (mean [SD]): 35.9 (13.3) years  Presence or absence of uncontrolled systemic hypertension (140/85mmHg) at the end of the trial: Not stated/unclear (initial blood pressure nifedipine, 154/60 (19/10)mmHg; initial blood pressure placebo: 155/62 (22/12)mmHg).		left ventricular overloading and, hence, the left ventricular end diastolic volume and mass in the population.
Scognamiglio 1994 <sup>146</sup>	Digoxin (n=74) Oral digoxin 0.25mg daily  Calcium channel blockers (n=69) Oral nifedipine 20mg twice daily  Concurrent medication/care: No additional information	Primary aortic regurgitation (N=143) Severity of disease: Severe Mechanism of disease: Rheumatic in 87/143. Aortic valve prolapse in 24/143. Bicuspid aortic valve in 32/143.  Defined by Doppler colour flow imaging.  Age (mean [SD]): 35.0 (13.0) years	All-cause mortality at 6 years Onset of symptoms or progression of NYHA class at 6 years Evidence of HVD progression on imaging (worsening of disease severity) at 6 years Need for heart valve intervention at 6 years Withdrawal due to adverse events at 6 years	Aim of the study: To determine whether this therapy delayed or reduced the need for aortic valve replacement.

		D		
		Presence or absence of uncontrolled systemic hypertension (140/85mmHg) at the end of the trial: Not stated/unclear (initial blood pressure digoxin: 150/58 (22/14), initial blood pressure nifedipine: 154/60 (20/8).)		
Stewart 2008 <sup>154</sup>	Diuretics (n=33) Oral eplerenone 50mg daily increased up to 100mg after one month if serum potassium, creatinine and systolic blood pressure were within normal limits and no adverse events.  Placebo (n=32) Oral placebo  Concurrent medication/care: Other medications were at the discretion of the patient's usual doctor.	Primary aortic [including bicuspid] stenosis (N=65) Severity of disease: Moderate to severe (peak velocity >3.0m/s)  Defined by Doppler ultrasound and echocardiography.  Age (mean [SD]): 67.5 (10.1) years Disease mechanism of aortic stenosis: Not stated/unclear Presence or absence of uncontrolled systemic hypertension (140/85mmHg) at the end of the trial: Not stated/unclear (initial blood pressure eplerenone, 145/83 (21/10)mmHg; initial blood pressure placebo: 144/81 (15/11)mmHg).	All-cause mortality at 19 months Cardiac mortality at 19 months Quality of life at 19 months Onset of symptoms or progression of NYHA class at 19 months Withdrawal due to adverse events at 19 months	Aim of the study: To determine whether the aldosterone-receptor antagonist eplerenone delays the onset of LV systolic dysfunction or reduces progression of LV hypertrophy assessed by cardiac magnetic resonance imaging in asymptomatic people with moderate to severe aortic stenosis. Additionally, to investigate the effects of eplerenone on non-invasive measures of LV diastolic function and progression of aortic valve stenosis.
Wisenbaugh 1994 <sup>177</sup>	Angiotensin-converting enzyme (ACE) inhibitors (n=13 AR, 14 MR)	Primary aortic regurgitation (n=23) Severity of disease: Severe	All-cause mortality at 6 months  Cardiac mortality at 6 months	Funding not stated Includes two strata, but outcomes reported separately.

Study	Intervention and comparison	Population	Outcomes	Comments
	Oral captopril 25mg three times a day for 6 months.	Mechanism of disease: Not stated	Onset of symptoms or progression of NYHA class at 6 months	Aim of the study: To investigate the effect of captopril on "remodelling" in the population.
	<b>Placebo</b> (n=10 AR,18 MR) Oral placebo	Defined by Doppler echocardiography and clinical examination.		
	Concurrent medication/care: Other vasodilating drugs were not used. People who were on furosemide were maintained on a constant dose.	Age (mean): 28.1 years Presence or absence of uncontrolled systemic hypertension (140/85mmHg) at the end of the trial: Not stated/unclear (initial blood pressure captopril, 131/46mmHg; initial blood pressure placebo: 144/57mmHg).		
		Primary mitral regurgitation (n=32) Severity of disease: Severe Mechanism of disease: Not stated		
		Defined by Doppler echocardiography and clinical examination.		
		Age (mean): 24.9 years Presence or absence of uncontrolled systemic hypertension (140/85mmHg) at the end of the trial: Not stated/unclear (initial blood		

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Study	Intervention and comparison	Population	Outcomes	Comments
		pressure captopril, 117/67mmHg; initial blood pressure placebo: 110/63mmHg).		

See Appendix D:for full evidence tables.

**2.1.2.3** Quality assessment of clinical studies included in the evidence review

Primary aortic [including bicuspid] stenosis 31.2.441

Table 4: Clinical evidence summary: ACE inhibitors compared to placebo

	No of			Anticipated absolute effects	
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with placebo	Risk difference with ACE inhibitors (95% CI)
All-cause mortality - not reported	-	-	Not estimable	-	-
Cardiac mortality - not reported	-	-	Not estimable	-	-
Health-related quality of life - not reported	-	-	Not estimable	-	-
Health-related quality of life - not reported	-	-	Not estimable	-	-
Onset of symptoms or progression in NYHA class - not reported	-	-	Not estimable	-	-
Evidence of HVD progression on imaging (worsening of disease severity) - not reported	-	-	Not estimable	-	-

	No of			Anticipated absolute effects	
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with placebo	Risk difference with ACE inhibitors (95% CI)
Need for heart valve intervention	83 (1 study) 12 months	⊕⊖⊖ VERY LOW <sub>1,2</sub> due to risk of bias, imprecision	RR 2.15 (0.42 to 11.1)	47 per 1000	54 more per 1000 (from 27 fewer to 475 more)
Exercise tolerance (change score) Exercise distance measured with treadmill exercise test	67 (1 study) 12 months	⊕⊕⊖ LOW <sub>1,3</sub> due to risk of bias		The mean exercise tolerance (change score) in the control groups was 29 meters	The mean exercise tolerance (change score) in the intervention groups was <b>49 meters lower</b> (61.59 to 36.41 lower)
Withdrawal due to adverse events	80 (1 study) 12 months	⊕⊖⊖ VERY LOW <sub>1,2</sub> due to risk of bias, imprecision	RR 2.21 (0.21 to 23.41)	24 per 1000	29 more per 1000 (from 19 fewer to 538 more)
1 Downgraded by 1 increment if	the majority of the ev	ridence was at high ris	sk of bias, and d	owngraded by 2 increments if t	he majority of the evidence was

<sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

3 MIDs used to assess imprecision were ±187.0

	No of Participants	evidence	Relative effect (95% CI)	Anticipated absolute effects	
Outcomes	(studies) Follow up			Risk with placebo	Risk difference with beta blockers (95% CI)
All-cause mortality - not reported	-	-	Not estimable	-	-
Cardiac mortality - not reported	-	-	Not estimable	-	-

<sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

Table 5: Clinical evidence summary: beta blockers compared to placebo

	No of Participants	Quality of the	Relative	Anticipated absolute effects	
Outcomes	(studies) Follow up	evidence (GRADE)	effect (95% CI)	Risk with placebo	Risk difference with beta blockers (95% CI)
Health-related quality of life (change score) Minnesota living with heart failure questionnaire. Scale from: 0 to 105 (high score is poor outcome).	38 (1 study) 5 months	⊕⊖⊖⊖ VERY LOW <sub>1,2,3</sub> due to risk of bias, imprecision		The mean health-related quality of life (change score) in the control groups was -1	The mean health-related quality of life (change score) in the intervention groups was <b>6 higher</b> (0.55 lower to 12.55 higher)
Health-related quality of life - not reported	-	-	Not estimable	-	
Onset of symptoms or progression in NYHA class (Copy) - not reported	-	-	Not estimable	-	-
Evidence of HVD progression on imaging (worsening of disease severity) - not reported	-	-	Not estimable	-	-
Need for heart valve intervention - not reported	-	-	Not estimable	-	-
Exercise tolerance (change score) 6-minute walk test distance	38 (1 study) 5 months	⊕⊖⊖ VERY LOW <sub>1,2,4</sub> due to risk of bias, imprecision		The mean exercise tolerance (change score) in the control groups was 14 meters	The mean exercise tolerance (change score) in the intervention groups was 12 meters lower (42.22 lower to 18.22 higher)
Withdrawal or dose reduction due to adverse events	38 (1 study) 5 months	⊕⊖⊖ VERY LOW <sub>1,2,5</sub> due to risk of bias, indirectness, imprecision	RR 2 (0.41 to 9.65)	105 per 1000	105 more per 1000 (from 62 fewer to 908 more)

<sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias
2 Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

(studies) evidence effect Risk difference with beta blockers (95% CI)  Outcomes (Studies) evidence (GRADE) (95% CI) Risk with placebo blockers (95% CI)		No of Participants	evidence	effect	Anticipated absolute effects	
	Outcomes	(studies)			Risk with placebo	

- 3 MIDs used to assess imprecision were ±5.0
- MIDs used to assess imprecision were ±21.0 5 Downgraded by 1 increment as the outcome includes people who had dose reductions or withdrawal due to adverse events

Table 6: Clinical evidence summary: diuretics compared to placebo

	No of Participants	Quality of the	Relative	Anticipated absolute effects		
Outcomes	(studies) Follow up	evidence (GRADE)	effect (95% CI)	Risk with placebo	Risk difference with diuretics (95% CI)	
All-cause mortality	61 (1 study) 19 months	⊕⊖⊖ VERY LOW <sub>1,2</sub> due to risk of bias, imprecision	RR 0.52 (0.05 to 5.4)	65 per 1000	31 fewer per 1000 (from 62 fewer to 286 more)	
Cardiac mortality	59 (1 study) 19 months	⊕⊖⊖ VERY LOW <sub>1,2</sub> due to risk of bias, imprecision	Peto OR 0.14 (0 to 7.06)	33 per 1000	<b>30 fewer per 1000</b> (from 120 fewer to 60 more) <sub>3</sub>	
Health-related quality of life - not reported	-	-	Not estimable	-	-	
Health-related quality of life (change score) SF-36 physical functioning subscale. Scale from: 0 to 100 (high score is good outcome).	59 (1 study) 12 months	⊕⊖⊖⊖ VERY LOW <sub>1,2,4</sub> due to risk of bias, imprecision		The mean health-related quality of life (change score) in the control groups was -9	The mean health-related quality of life (change score) in the intervention groups was <b>4 higher</b> (6.5 lower to 14.5 higher)	
Health-related quality of life (change score) SF-36 role physical subscale.	59 (1 study) 12 months	⊕⊖⊖⊖ VERY LOW <sub>1,2,4</sub>		The mean health-related quality of life (change score)	The mean health-related quality of life (change score) in the intervention groups was	

	No of Participants	Quality of the	Relative	Anticipated absolute effects		
Outcomes	(studies) evidence effect Follow up (GRADE) (95% CI)		Risk with placebo	Risk difference with diuretics (95% CI)		
Scale from: 0 to 100 (high score is good outcome).		due to risk of bias, imprecision		in the control groups was -12	3 higher (15.12 lower to 21.12 higher)	
Onset of symptoms or progression of NYHA class	59 (1 study) 19 months	⊕⊖⊖ VERY LOW <sub>1,2</sub> due to risk of bias, imprecision	RR 1.34 (0.7 to 2.57)	333 per 1000	<b>113 more per 1000</b> (from 100 fewer to 523 more)	
Evidence of HVD progression on imaging (worsening of disease severity) - not reported	-	-	Not estimable	-	-	
Need for heart valve intervention - not reported	-	-	Not estimable	-	-	
Withdrawal due to adverse events	62 (1 study) 19 months	⊕⊖⊖ VERY LOW <sub>1,2</sub> due to risk of bias, imprecision	Peto OR 6.94 (0.14 to 350.54)	0 per 1000	<b>30 more per 1000</b> (from 50 fewer to 120 more) <sub>3</sub>	

<sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

#### Table 7: Clinical evidence summary: statins compared to placebo

		Quality of		Anticipated absolute effects		
Outcomes	No of Participants (studies) Follow up	the evidence (GRADE)	Relative effect (95% CI)	Risk with placebo	Risk difference with statins (95% CI)	

<sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

<sup>3</sup> Absolute effect calculated manually using risk difference as zero events in one arm of the study

 $_{\rm 4}$  MIDs used to assess imprecision were  $\pm 3.0$ 

		Quality of		Anticipated absolute effects	
Outcomes	No of Participants (studies) Follow up	the evidence (GRADE)	Relative effect (95% CI)	Risk with placebo	Risk difference with statins (95% CI)
All-cause mortality	2189 (3 studies) 4.3 years	⊕⊖⊖ VERY LOW <sub>1,2,3</sub> due to risk of bias, indirectness, imprecision	RR 1.01 (0.79 to 1.3)	44 per 1000	0 more per 1000 (from 9 fewer to 13 more)
All-cause mortality (time to event)	1873 (1 study) 4.4 years	⊕⊖⊖ VERY LOW <sub>1,2,3</sub> due to risk of bias, indirectness, imprecision	HR 1.04 (0.79 to 1.37)	108 per 1000	4 more per 1000 (from 22 fewer to 37 more)
Cardiac mortality	2344 (4 studies) 3.7 years	⊕⊕⊖ LOW <sub>2,3</sub> due to indirectness, imprecision	RR 0.75 (0.54 to 1.03)	52 per 1000	13 fewer per 1000 (from 24 fewer to 2 more)
Cardiac mortality (time to event)	1873 (1 study) 4.4 years	⊕⊖⊖ VERY LOW <sub>1,2,3</sub> due to risk of bias, indirectness, imprecision	HR 0.83 (0.56 to 1.23)	60 per 1000	10 fewer per 1000 (from 26 fewer to 13 more)
Health-related quality of life - not reported	-	-	Not estimable	-	-
Health-related quality of life - not reported	-	-	Not estimable	-	-

		Quality of		Anticipated absolute effects	
Outcomes	No of Participants (studies) Follow up	the evidence (GRADE)	Relative effect (95% CI)	Risk with placebo	Risk difference with statins (95% CI)
Onset of symptoms or progression of NYHA class	2028 (2 studies) 3.2 years	⊕⊖⊖ VERY LOW <sub>2,3</sub> due to indirectness, imprecision	RR 0.99 (0.59 to 1.66)	44 per 1000	0 fewer per 1000 (from 18 fewer to 29 more)
Onset of symptoms or progression of NYHA class (time to event)	1873 (1 study) 4.4 years	⊕⊖⊖ VERY LOW <sub>1,2,3</sub> due to risk of bias, indirectness, imprecision	HR 1.09 (0.62 to 1.92)	25 per 1000	2 more per 1000 (from 9 fewer to 22 more)
Evidence of HVD progression on imaging (worsening of disease severity) - not reported	-	-	Not estimable	-	-
Need for heart valve intervention	2346 (4 studies) 3.7 years	⊕⊖⊖ VERY LOW <sub>2,3,4</sub> due to inconsistency , indirectness, imprecision	RR 0.93 (0.7 to 1.24)	222 per 1000	16 fewer per 1000 (from 67 fewer to 53 more)
Need for heart valve intervention (time to event)	1873 (1 study) 4.4 years	⊕⊕⊖⊖ LOW <sub>1,2</sub> due to risk of bias, indirectness	HR 1 (0.84 to 1.19)	299 per 1000	0 fewer per 1000 (from 41 fewer to 46 more)
Withdrawal due to adverse events	48 (1 study) 6 months	⊕⊖⊝⊖ VERY LOW <sub>1,3</sub>	Peto OR 6.82	0 per 1000	<b>40 more per 1000</b> (from 70 fewer to 150 more) <sub>5</sub>

		Quality of		Anticipated absolute effects		
No of Participants the (studies) evidence utcomes Follow up (GRADE)	Relative effect (95% CI)	Risk with placebo	Risk difference with statins (95% CI)			
		due to risk of bias, imprecision	(0.13 to 344.93)			
Withdrawal due to adverse events	2296 (3 studies) 3.3 years	⊕⊕⊖ LOW <sub>2,3</sub> due to indirectness, imprecision	RR 1.15 (0.94 to 1.4)	131 per 1000	20 more per 1000 (from 8 fewer to 52 more)	

<sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

#### Primary aortic regurgitation

Table 8: Clinical evidence summary: ACE inhibitors compared to placebo/no treatment

	No of			Anticipated absolute effects	
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with Placebo/no treatment	Risk difference with ACE-inhibitors (95% CI)
All-cause mortality	63 (1 study) 7 years	⊕⊖⊖ VERY LOW <sub>1,2</sub> due to risk of bias, imprecision	RR 0.97 (0.06 to 14.82)	32 per 1000	1 fewer per 1000 (from 30 fewer to 442 more)

<sup>2</sup> Downgraded by 1 increment as one study included a statin and ezetimibe in the intervention group

<sup>3</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

<sup>4</sup> Downgraded by 1 as the point estimate varies widely across studies, with subgroup analysis not being possible due to the difference being seen in one study

<sup>5</sup> Absolute effect calculated manually using risk difference as zero events in one arm of the study

	No of			Anticipated absolute effects	
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with Placebo/no treatment	Risk difference with ACE-inhibitors (95% CI)
Cardiac mortality	63 (1 study) 7 years	⊕⊖⊖ VERY LOW <sub>1,2</sub> due to risk of bias, imprecision	Peto OR 0.13 (0 to 6.61)	32 per 1000	<b>30 fewer per 1000</b> (from 120 fewer to 50 more) <sub>3</sub>
Health-related quality of life - not reported	-	-	Not estimable	-	-
Health-related quality of life - not reported	-	-	Not estimable	-	-
Onset of symptoms or progression of NYHA class	83 (2 studies) 7 years	⊕⊕⊖ LOW <sub>1,5</sub> due to risk of bias, inconsistency	RD 0 (-0.13 to 0.22)	200 per 1000	<b>40 more per 1000</b> (from 130 fewer to 220 more) <sub>4</sub>
Evidence of HVD progression on imaging (worsening of disease severity)	63 (1 study) 7 years	⊕⊖⊖ VERY LOW <sub>1,2</sub> due to risk of bias, imprecision	RR 1.36 (0.71 to 2.58)	323 per 1000	<b>116 more per 1000</b> (from 94 fewer to 510 more)
Need for heart valve intervention	63 (1 study) 7 years	⊕⊖⊖ VERY LOW <sub>1,2</sub> due to risk of bias, imprecision	RR 1.29 (0.74 to 2.27)	387 per 1000	<b>112 more per 1000</b> (from 101 fewer to 491 more)
Withdrawal due to adverse events	63 (1 study) 7 years	⊕⊖⊖ VERY LOW <sub>1,2</sub> due to risk of bias, imprecision	Peto OR 7.65 (0.77 to 76.34)	0 per 1000	<b>90 more per 1000</b> (from 20 fewer to 210 more) <sub>3</sub>

<sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

<sup>&</sup>lt;sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

<sup>3</sup> Absolute effect calculated manually using risk difference as zero events in one arm of the study

<sup>4</sup> Absolute effect calculated manually using risk difference as zero events in both arms of a study

<sup>5</sup> Downgraded by 1 increment as zero events in both arms of one study

2 Table 9: Clinical evidence summary: ACE inhibitors compared to calcium channel blockers

				Anticipated abso	ute effects
Outcomes	No of Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with calcium channel blockers	Risk difference with ACE inhibitors (95% CI)
All-cause mortality	64 (1 study) 7 years	⊕⊖⊖ VERY LOW₁ due to risk of bias, imprecision	RR 1 (0.07 to 15.3)	31 per 1000	<b>0 fewer per 1000</b> (from 29 fewer to 447 more)
Cardiac mortality	64 (1 study) 7 years	⊕⊖⊖ VERY LOW <sub>1,3</sub> due to risk of bias, imprecision	Peto OR 0.14 (0 to 6.82)	31 per 1000	<b>30 fewer per 1000</b> (from 110 fewer to 50 more) <sub>2</sub>
Health-related quality of life - not reported	-	-	Not estimable	-	-
Health-related quality of life - not reported	-	-	Not estimable	-	-
Onset of symptoms or progression of NYHA class	89 (2 studies) 4.8 years	⊕⊖⊖ VERY LOW₃ due to risk of bias, inconsistency	-RD 0.04 (-0.12 to 0.21)	125 per 1000	<b>40 more per 1000</b> (from 120 fewer to 210 more) <sub>4</sub>
Evidence of HVD progression on imaging (worsening of disease severity)	89 (2 studies) 4.8 years	⊕⊖⊖ VERY LOW <sub>1,3,6</sub> due to risk of bias, inconsistency, imprecision	RR 1.15 (0.62 to 2.13)	240 per 1000	<b>40 more per 1000</b> (from 140 fewer to 230 more)₅
Need for heart valve intervention	64 (1 study) 7 years	⊕⊖⊖ VERY LOW <sub>1,3</sub> due to risk of bias, imprecision	RR 1.23 (0.71 to 2.12)	406 per 1000	<b>93 more per 1000</b> (from 118 fewer to 455 more)

				Anticipated absolute effects		
Outcomes	No of Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with calcium channel blockers	Risk difference with ACE inhibitors (95% CI)	
Withdrawal due to adverse events	64 (1 study) 7 years	⊕⊖⊖ VERY LOW <sub>1,3</sub> due to risk of bias, imprecision	RR 0.43 (0.12 to 1.51)	219 per 1000	<b>125 fewer per 1000</b> (from 193 fewer to 112 more)	

- 1 Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs
- 2 Absolute effect calculated manually using risk difference as zero events in one arm of the study
- 3 Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias
- 4 Absolute effect calculated manually using risk difference as zero events in both arms of one study
- 5 Absolute effect calculated manually using risk difference as zero events in one arm of a study
- 6 Downgraded by 1 increment as zero events in one of the studies included

#### Table 10: Clinical evidence summary: ARBs compared to beta blockers

	No of			Anticipated absolute effects		
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with beta blockers	Risk difference with ARBs (95% CI)	
All-cause mortality - not reported	-	-	Not estimable	-	-	
Cardiac mortality - not reported	-	-	Not estimable	-	-	
Health-related quality of life - not reported	-	-	Not estimable	-	-	
Health-related quality of life - not reported	-	-	Not estimable	-	-	
Onset of symptoms or progression in NYHA class - not reported	-	-	Not estimable	-	-	

Table 11: Clinical evidence summary: Beta blockers compared to placebo

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No of				Anticipated absolute effects					
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with placebo	Risk difference with beta blockers (95% CI)				
All-cause mortality - not reported	-	-	Not estimable	-	-				

	No of			Anticipated absolute effect	cts	
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with beta blockers	Risk difference with ARBs (95% CI)	
Evidence of HVD progression on imaging (worsening of disease severity) - not reported	-	-	Not estimable	-	-	
Need for heart valve intervention - not reported	-	-	Not estimable	-	-	
Exercise tolerance (final value) exercise work rate using an ergometer	34 (1 study) 3 weeks	⊕⊖⊖ VERY LOW <sub>1,2,3,4</sub> due to risk of bias, indirectness, imprecision		The mean exercise tolerance (final value) in the control groups was 29 Watts	The mean exercise tolerance (final value) in the intervention groups was <b>0 Watts higher</b> (4.75 lower to 4.75 higher) <sup>5</sup>	

<sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

<sup>2</sup> Downgraded by 1 increment as follow up less than 1 month

<sup>3</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

<sup>4</sup> MIDs used to assess imprecision were ±4.0

<sup>5</sup> Insufficient information available to conduct a paired analysis

	No of			Anticipated absolute effe	cts
Outcomes	Participants Quality of the Relative (studies) evidence effect Follow up (GRADE) (95% CI)			Risk with placebo	Risk difference with beta blockers (95% CI)
Cardiac mortality - not reported	-	-	Not estimable	-	-
Onset of symptoms or progression in NYHA class - not reported	-	-	Not estimable	-	-
Quality of life (final value) EuroQol visual analogue scale. Scale from: 0 to 100 (high score is good outcome).	72 (1 study) 6 months	⊕⊕⊖ LOW <sub>1,2,3</sub> due to risk of bias, imprecision		The mean quality of life (final value) in the control groups was 82	The mean quality of life (final value) in the intervention groups was  3 higher (2.7 lower to 8.7 higher)
Quality of life (final value) KCCQ. Scale from: 0 to 100 (high score is good outcome).	72 (1 study) 6 months	⊕⊕⊖ LOW <sub>1,2,4</sub> due to risk of bias, imprecision		The mean quality of life (final value) in the control groups was 96	The mean quality of life (final value) in the intervention groups was <b>2 higher</b> (17.76 lower to 21.76 higher)
Evidence of HVD progression on imaging (worsening of disease severity) - not reported	-	-	Not estimable	-	-
Need for heart valve intervention - not reported	-	-	Not estimable	-	-
Exercise tolerance Peak work (bicycle ergometer) (high score if good outcome)	72 (1 study) 6 months)	⊕⊕⊖⊖ LOW <sub>1,2,5</sub> due to risk of bias, imprecision		The mean exercise tolerance in the control groups was 241 watts	The mean exercise tolerance in the intervention groups was  12 watts lower  (40.64 lower to 16.64 higher)

<sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

<sup>&</sup>lt;sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

 $_3$  MIDs used to assess imprecision were  $\pm 5.00$ 

<sup>4</sup> MIDs used to assess imprecision were ±21.39

<sup>&</sup>lt;sub>5</sub> MIDs used to assess imprecision were ±31.50

# 1 Table 12: Clinical evidence summary: Calcium channel blockers compared to placebo/no treatment

	No of			Anticipated absolute effects		
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with Placebo/no treatment	Risk difference with Calcium channel blockers (95% CI)	
All-cause mortality	63 (1 study) 7 years	⊕⊖⊖ VERY LOW <sub>1,2</sub> due to risk of bias, imprecision	RR 0.97 (0.06 to 14.82)	32 per 1000	1 fewer per 1000 (from 30 fewer to 442 more)	
Cardiac mortality	64 (1 study) 7 years	⊕⊖⊖ VERY LOW <sub>1,2</sub> due to risk of bias, imprecision	RR 1 (0.07 to 15.3)	31 per 1000	0 fewer per 1000 (from 29 fewer to 443 more)	
Health-related quality of life - not reported	-	-	Not estimable	-	-	
Health-related quality of life - not reported	-	-	Not estimable	-	-	
Onset of symptoms or progression of NYHA class	63 (1 study) 7 years	⊕⊖⊖ VERY LOW <sub>1,2</sub> due to risk of bias, imprecision	RR 0.97 (0.42 to 2.26)	258 per 1000	8 fewer per 1000 (from 150 fewer to 325 more)	
Evidence of HVD progression on imaging (worsening of disease severity)	63 (1 study) 7 years	⊕⊖⊖ VERY LOW <sub>1,2</sub> due to risk of bias, imprecision	RR 0.97 (0.47 to 2)	323 per 1000	10 fewer per 1000 (from 171 fewer to 323 more)	
Need for heart valve intervention	135 (2 studies) 7 years	⊕⊖⊖ VERY LOW <sub>1,4,5</sub> due to risk of bias, inconsistency, imprecision	RD 0.01 (-0.11 to 0.13)	179 per 1000	<b>10 more per 1000</b> (from 110 fewer to 130 more) <sub>3</sub>	
Withdrawal due to adverse events	135 (2 studies) 7 years	⊕⊖⊖ VERY LOW <sub>1,2,4</sub> due to risk of bias, inconsistency, imprecision	RR 8.51 (1.12 to 64.44)	0 per 1000	<b>120 more per 1000</b> (from 30 more to 200 more) <sub>6</sub>	

	No of			Anticipated abs	solute effects
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with Placebo/no treatment	Risk difference with Calcium channel blockers (95% CI)

- 1 Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias
- 2 Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs
- 3 Absolute effect calculated manually using risk difference as zero events in both arms of a study
- 4 Downgraded by 1 increment as zero events in both arms of one study
- 5 Downgraded by 2 increments as zero events in both arms of a study and OIS <80%
- 6 Absolute effect calculated manually using risk difference as zero events one arm of the study

# Table 13: Clinical evidence summary: Digoxin compared to calcium channel blockers

				Anticipated absolu	te effects
Outcomes	No of Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with Calcium channel blockers	Risk difference with Digoxin (95% CI)
All-cause mortality	135 (1 study) 6 years	⊕⊖⊖ VERY LOW <sub>2,3</sub> due to risk of bias, imprecision	Peto OR 6.88 (0.14 to 347.65)	0 per 1000	<b>10 more per 1000</b> (from 30 fewer to 50 more) <sub>1</sub>
Cardiac mortality - not reported	-	-	Not estimable	-	-
Health-related quality of life - not reported	-	-	Not estimable	-	-
Health-related quality of life - not reported	-	-	Not estimable	-	-
Onset of symptoms of progression of NYHA class	135 (1 study) 6 years	⊕⊕⊝ LOW <sub>2,3</sub> due to risk of bias, imprecision	RR 2.63 (1.11 to 6.26)	92 per 1000	<b>150 more per 1000</b> (from 10 more to 486 more)
Evidence of HVD progression on imaging (worsening of disease severity)	135 (1 study) 6 years	⊕⊕⊖⊖ LOW <sub>2,3</sub> due to risk of bias, imprecision	Peto OR 7.30 (1.23 to 43.33)	0 per 1000	<b>70 fewer per 1000</b> (from 10 more to 140 more) <sub>1</sub>

			Anticipated absolute effects		
Outcomes	No of Participants (studies) Follow up	es) evidence		Risk with Calcium channel blockers	Risk difference with Digoxin (95% CI)
Need for heart valve intervention	135 (1 study) 6 years	⊕⊕⊖ LOW <sub>2,3</sub> due to risk of bias, imprecision	RR 3.10 (1.33 to 7.22)	92 per 1000	<b>194 more per 1000</b> (from 30 more to 574 more)
Withdrawal due to adverse events	135 (1 study) 6 years	⊕⊕⊖⊖ LOW <sub>2,5</sub> due to risk of bias, imprecision	RD 0 (-0.03 to 0.03)	0 per 1000	<b>0 fewer per 1000</b> (from 30 fewer to 30 more) <sub>4</sub>

<sup>&</sup>lt;sup>1</sup> Absolute effect calculated from risk difference due to zero events in one study arm

# 2.1.2.3.1 Primary mitral stenosis

3 No studies identified.

# 4 2.1.2.3.2 Primary mitral regurgitation

# 5 Table 14: Clinical evidence summary: ACE inhibitors compared to placebo

	No of			Anticipated absolute effects		
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with placebo	Risk difference with ACE inhibitors (95% CI)	
All-cause mortality	45 (2 studies) 6-12 months	⊕⊖⊖ VERY LOW <sub>2,3,4,5</sub> due to risk of bias,	-RD -0.04 (- 0.18 to 0.11)	29 per 1000	<b>40 fewer per 1000</b> (from 180 fewer to 110 more) <sub>1</sub>	

<sup>&</sup>lt;sup>2</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

<sup>3</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

<sup>4</sup> Absolute effect calculated from risk difference due to zero events in both study arms

<sup>5</sup> Downgraded by 1 increment as sample size is between 75 and 350 with zero events in both arms

	No of			Anticipated absolute effe	ects
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with placebo	Risk difference with ACE inhibitors (95% CI)
		inconsistency, indirectness, imprecision			
Cardiac mortality	45 (2 studies) 6-12 months	⊕⊖⊖ VERY LOW <sub>2,3,4,5</sub> due to risk of bias, inconsistency, indirectness, imprecision	-RD -0.04 (- 0.18 to 0.11)	29 per 1000	<b>40 fewer per 1000</b> (from 180 fewer to 110 more) <sub>1</sub>
Quality of life (change score) Life quality index. Scale from: 1 to 6 (high score is good outcome).	16 (1 study) 6 months	⊕⊖⊖ VERY LOW <sub>2,3,6</sub> due to risk of bias, indirectness		The mean quality of life (change score) in the control groups was 0.4	The mean quality of life (change score) in the intervention groups was <b>0.2 lower</b> (1.03 lower to 0.63 higher)
Quality of life (change score) Life quality index. Scale from: 1 to 6 (high score is good outcome).	16 (1 study) 1 years	⊕⊖⊖ VERY LOW <sub>2,3,6</sub> due to risk of bias, indirectness		The mean quality of life (change score) in the control groups was 0.4	The mean quality of life (change score) in the intervention groups was <b>0.1 lower</b> (0.93 lower to 0.73 higher)
Onset of symptoms or progression of NYHA class	77 (2 studies) 6-12 months	⊕⊖⊖ VERY LOW <sub>2,3,7</sub> due to risk of bias, indirectness, imprecision	RR 0.17 (0.02 to 1.26)	120 per 1000	<b>140 fewer per 1000</b> (from 270 fewer to 10 fewer) <sub>1</sub>
Evidence of HVD progression on imaging (worsening of disease severity) - not reported	-	-	Not estimable	-	-
Need for heart valve intervention	48 (1 study) 1 years	⊕⊖⊖ VERY LOW <sub>2,7</sub>	Peto OR 0.11 (0 to 5.76)	46 per 1000	<b>50 fewer per 1000</b> (from 160 fewer to 70 more) <sub>1</sub>

	No of			Anticipated absolute effe	cts
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with placebo	Risk difference with ACE inhibitors (95% CI)
		due to risk of bias, imprecision			
Exercise tolerance (change score) Bruce Protocol treadmill exercise time (seconds)	16 (1 study) 1 years	⊕⊖⊖ VERY LOW <sub>2,3,7,8</sub> due to risk of bias, indirectness, imprecision		The mean exercise tolerance (change score) in the control groups was 18 seconds	The mean exercise tolerance (change score) in the intervention groups was <b>21 seconds higher</b> (42.97 lower to 84.97 higher)
Exercise tolerance (final value) oxygen uptake at peak exercise (mL/min)	47 (1 study) 1 years	⊕⊖⊖ VERY LOW <sub>2,7,9</sub> due to risk of bias, imprecision		The mean exercise tolerance (final value) in the control groups was 1433 mL/min	The mean exercise tolerance (final value) in the intervention groups was <b>361 mL/min higher</b> (50.91 to 671.09 higher)
Withdrawal due to adverse events	21 (1 study) 1 years	⊕⊖⊖ VERY LOW <sub>2,3,7</sub> due to risk of bias, indirectness, imprecision	RR 4.4 (0.59 to 33.07)	91 per 1000	<b>309 more per 1000</b> (from 37 fewer to 1000 more)

- <sub>1</sub> Absolute effect calculated manually using risk difference as zero events in the studies
- <sup>2</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias
- 3 Downgraded by 1 increment as some of the participants in one study may have had congenital valvular heart disease
- 4 Downgraded by 1 increment as one study has zero events in both arms, and one has zero events in one arm
- 5 Downgraded by 2 increments as calculated power was less than 80%
- $_{\rm 6}$  MIDs used to assess imprecision were  $\pm 1.12$
- 7 Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs
- 8 MIDs used to assess imprecision were ±66.90
- 9 MIDs used to assess imprecision were ±270.50

# Table 15: Clinical evidence summary: beta blockers compared to placebo

				Anticipate	d absolute effects
Outcomes	No of Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with placebo	Risk difference with beta blockers (95% CI)
All-cause mortality	37 (1 study) 2 years	⊕⊖⊖ VERY LOW <sub>2,3</sub> due to risk of bias, imprecision	OR 7.01 (0.14 to 353.8)	0 per 1000	<b>50 more per 1000</b> (from 80 fewer to 190 more) <sub>1</sub>
Cardiac mortality	37 (1 study) 2 years	⊕⊖⊖ VERY LOW <sub>2,3</sub> due to risk of bias, imprecision	OR 7.01 (0.14 to 353.8)	0 per 1000	<b>50 more per 1000</b> (from 80 fewer to 190 more) <sub>1</sub>
Health-related quality of life - not reported	-	-	Not estimable	-	-
Health-related quality of life - not reported	-	-	Not estimable	-	-
Onset of symptoms or progression in NYHA class - not reported	-	-	Not estimable	-	-
Evidence of HVD progression on imaging (worsening of disease severity) - not reported	-	-	Not estimable	-	-
Need for heart valve intervention	36 (1 study) 2 years	⊕⊖⊖ VERY LOW <sub>2,3</sub> due to risk of bias, imprecision	RR 0.33 (0.08 to 1.44)	333 per 1000	223 fewer per 1000 (from 306 fewer to 147 more)
Serious adverse events	36 (1 study) 2 years	⊕⊖⊖⊖ VERY LOW <sub>2,3,4</sub> due to risk of bias, indirectness, imprecision	RR 0.43 (0.13 to 1.4)	389 per 1000	222 fewer per 1000 (from 338 fewer to 156 more)

Absolute effect calculated manually using risk difference as zero events in the studies
 Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

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				Anticipated absolute effects		
Outcomes	No of Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with placebo	Risk difference with beta blockers (95% CI)	
<ul> <li>Downgraded by 1 increment if the confidence interval cro</li> <li>Downgraded by 1 increment as the study does not report</li> </ul>	•		ice interval cros	ssed both MI	Ds	

- 2 2.1.2.3.3 Primary tricuspid regurgitation
- 3 No studies identified.
- 4 2.1.2.3.4 Secondary valvular heart disease mitral regurgitation and tricuspid regurgitation
- 5 No studies identified.
- 7 See Appendix F: for full GRADE tables.

# 1 2.1.3 Economic evidence

## 2 2.1.3.1 Included studies

3 No health economic studies were included.

## 4 2.1.3.2 Excluded studies

- 5 No relevant health economic studies were excluded due to assessment of limited
- 6 applicability or methodological limitations.
- 7 See also the health economic study selection flow chart in Appendix G:.

# 8 2.1.3.3 Health economic modelling

9 This area was not prioritised for new cost-effectiveness analysis.

## 10 **2.1.3.4 Unit costs**

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- 11 The following relevant unit costs have been included to inform the committee of the cost
- 12 implications of different pharmacological management strategies.

Table 16. Unit costs for different drugs used for pharmacological management of people with heart failure and concomitant heart valve disease

Class	Drug	Dose (tablet unless specified)	Ur	nit cost
Alpha blockers	doxasozin	2mg	£	0.04
ACE inhibitors	ramipril	1.25mg	£	0.07
		2.5mg	£	0.15
		5mg	£	0.17
		10mg	£	0.18
	captopril	12.5mg	£	0.02
		25mg	£	0.01
		50mg	£	0.03
	enalapril	2.5mg	£	0.18
		5mg, 10mg, 20mg	£	0.06
	lisinopril	2.5mg, 5mg, 10mg, 20mg	£	0.03
	quinapril	2.5mg, 5mg, 10mg	£	0.31
		20mg	£	0.39
		40mg	£	0.13
	fosinopril	10mg	£	0.15
		20mg	£	0.14
Angiotensin II	candesartan cilexitil	2mg	£	0.22
receptor blockers (ARBs)		4mg	£	80.0
2.23(0.0 (7.11(120)		8mg	£	0.04
		16mg	£	0.06
		32mg	£	0.06

	losartan	12.5mg	£	0.11
		25mg	£	0.12
		50mg	£	0.07
		100mg	£	0.07
Beta blockers	bisoprolol	1.25mg	£	0.03
		3.75mg	£	0.03
		5mg	£	0.02
		10mg	£	0.03
	carvedilol	3.125mg	£	0.03
		6.25mg	£	0.03
		12.5mg	£	0.03
		25mg	£	0.04
	nebivolol	2.5mg	£	0.42
		5mg	£	0.16
		10mg	£	0.92
Diuretics	furosemide	20mg tablet	£	0.05
		40mg tablet	£	0.07
		10 mg per 1 ml solution for injection	£	1.74
	bumetanide	1mg tablet	£	0.05
		5mg tablet	£	0.25
	torasemide	2.5mg tablet	£	0.14
		5mg	£	0.20
		10mg	£	0.29
Calcium channel blockers	amlopodine	5mg, 10mg	£	0.03
Digoxin	-	62.5 micrograms	£	0.05
		125 micrograms	£	0.05
Nitrates	Isosorbide dinitrate	10mg	£	0.24
	Nitroprusside		No tariff price	e available
Statins	Atorvastatin	10mg	£	0.03
		20mg	£	0.03
		80mg	£	0.07
	Fluvastin	20mg	£	0.08
		40mg	£	0.09
		80mg (modified release capsule)	£	0.69
	Pravastatin	10mg	£	0.03
		20mg	£	0.04
		40mg	£	0.05
	Rosuvastatin	5mg	£	0.06
		10mg	£	0.05

		20mg	£	0.07
		40mg	£	0.10
		10mg	£	0.03
		20mg	£	0.03
		40mg	£	0.04
		80mg	£	0.06

1 Source: BNF 2018<sup>79</sup>

# 2 **2.1.4 Evidence statements**

# 3 2.1.4.1 Clinical evidence statements

- 4 See the summary of evidence in Table, Table, Table, Table, Table, Table, Table,
- 5 Table, Table, Table and Table 15.

# 6 2.1.4.2 Health economic evidence statements

8 No relevant economic evaluations were identified.

## 9 2.1.5 The committee's discussion of the evidence

# 10 **2.1.6 Interpreting the evidence**

# 11 2.1.6.1 The outcomes that matter most

- 12 The critical outcomes were all-cause mortality, cardiac mortality, health-related quality of life,
- onset of symptoms or progression in NYHA class, evidence of HVD progression on imaging
- 14 and need for heart valve intervention. The important outcomes were exercise tolerance and
- withdrawal from the trial due to adverse events. Exercise tolerance was considered important
- due to its impact on quality of life and as a measure of symptom burden and withdrawal due
- 17 to adverse events would provide information on any severe events associated with any of the
- 18 drugs.

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- 19 Physiological outcomes were not included as they are not clinically relevant endpoints, and
- the outcomes they aim to predict that are important to patients are captured by the included
- 21 outcomes.
- 22 There was very limited evidence. All outcomes were reported in at least one study. However,
- 23 there were gaps in outcomes reported for specific strata. For the aortic stenosis and mitral
- 24 regurgitation strata, no studies reported evidence of HVD progression on imaging. For the
- 25 aortic regurgitation stratum, no studies reported health-related quality of life at ≥12 months or
- 26 withdrawal due to adverse events at <6 months.

# 27 **2.1.6.2** The quality of the evidence

- No relevant RCTs for mitral stenosis, tricuspid regurgitation and secondary heart valve
- 29 disease were identified. No relevant RCTs investigating the use of alpha blockers or nitrates
- 30 were identified. Seventeen RCTs were included in this review and evidence was only
- 31 available for the following comparisons:
- 32 Aortic stenosis
- 33 o ACE-I versus placebo
- o Beta-blocker versus placebo

- 1 o Diuretic versus placebo
- 2 o Statin versus placebo
- Aortic regurgitation
- 4 o ACE-I versus placebo/no treatment
- 5 o ACE-I versus calcium channel blocker
- 6 o ARB versus beta-blocker
- 7 o Beta-blocker versus placebo
- 8 o Calcium channel blocker versus placebo/no treatment
- 9 o Digoxin versus calcium channel blocker
- 10 Mitral regurgitation

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- ACE-I versus placebo
- o Beta-blocker versus placebo

13 Evidence ranged from low to very low quality, with the majority of the evidence being of very 14 low quality. Evidence was mainly downgraded due to risk of bias and imprecision. Analyses 15 frequently included only a small number of participants and had low event rates resulting in 16 great uncertainty. Additionally, some evidence was considered to be indirect because of 17 inclusion of additional pharmacological agents not stated in the protocol (for example, ezetimibe with statins) or inclusion of people with congenital valve disease (while including 18 19 an adequate proportion of the population without congenital valve disease to still fulfil the protocolised inclusion criteria). One outcome (withdrawal due to adverse events at 1-7 years, 20 for the comparison between calcium channel blockers and no treatment in primary aortic 21 22 regurgitation) showed inconsistency with heterogeneity that could not be explained by

## 24 2.1.6.3 Benefits and harms

subgroup analysis.

# 2.1.6.3.1 Aortic stenosis

The evidence showed a small clinically important benefit of statins for cardiac mortality, with no clinically important difference in all-cause mortality, onset of symptoms, need for heart valve intervention, and withdrawal due to adverse events; however, for all of these outcomes confidence intervals demonstrated uncertainty in the effect. Although the evidence from these studies suggested increased withdrawal due to adverse events in the statin group compared to placebo, there was also uncertainty in this effect and the absolute effect was not considered to represent a clinically important difference. The committee agreed that statins are unlikely to directly affect the severity of the aortic valve lesion, which possibly explains why there was no clinically important difference in need for heart valve intervention observed, but they may help with other confounding variables that we cannot determine from this evidence that influence cardiac mortality. For example, aortic stenosis may be a marker for increased cardiovascular risk. Two of the four studies that were included in this analysis excluded people with a history of arteriopathy (including coronary artery disease, cerebrovascular disease, and peripheral vascular disease), although no history of arteriopathy does not necessarily mean it is not currently present. Both of these studies showed a clinically important benefit for cardiac mortality. It was agreed that the relative effect size showing a 25% reduced chance of death from cardiac causes, relating to 13 per 1000 fewer cases, would be important to people with aortic stenosis and that the metaanalysis was of sufficient size to provide evidence for this. Despite the evidence being graded low to very low quality for this comparison and uncertainty identified for all outcomes, the committee agreed that due to the likely impact on general cardiovascular health a crossreference to the NICE guideline on lipid modification was appropriate, which includes recommendations on statin use. . The committee highlighted that although this review

- 1 focused on those without heart failure and the pharmacological review protocol on heart
- 2 valve disease with heart failure did not include statins, the results could also be applied to
- 3 those with aortic stenosis and heart failure as statins are thought to affect general
- 4 cardiovascular health rather than having an effect on aortic stenosis itself. Statins were not
- 5 included in the heart valve disease with heart failure review protocol as this review aimed to
- 6 focus on drugs that are commonly used to treat heart failure, which does not include statins.
- 7 The committee noted that statin use in moderate-to-severe aortic stenosis may be too late,
- 8 and that advocates for the use of statins in aortic stenosis believe that starting statins in
- 9 people with mild aortic stenosis may have more benefit in preventing progression of heart
- 10 valve disease. This population was excluded in this review as it was noted that mild valve
- 11 disease is rarely followed up and rarely progresses, and recommendations could therefore
- 12 not be made.
- 13 There was insufficient evidence to draw conclusions about the relative benefits and harms of
- ACE-I, beta-blockers and diuretics based on the evidence available. Although ACE-I and
- 15 beta-blockers showed increased events in those with moderate or severe aortic stenosis in
- 16 terms of need for heart valve intervention at 12 months and withdrawal due to adverse
- events, respectively, in both cases only one small study with evidence graded very low
- 18 quality was available for each comparison and imprecise estimates were reported that did
- 19 not show a large enough difference in effect for the committee to be confident in the findings,
- 20 with a difference of only two events between the groups in both cases that could have
- 21 occurred by chance. Conversely, possible benefits in outcomes were seen when diuretics
- were compared to placebo. This was in one small study, with evidence graded very low
- 23 quality and uncertainty observed for all outcomes, where the pharmacological agent was
- 24 eplerenone, a mineralocorticoid receptor antagonist. The committee agreed that this may not
- 25 be representative of other diuretics that have a different mechanism of action (as eplerenone
- acts on extrarenal pathways). In addition, there was a difference of only 1-3 events across
- 27 the dichotomous outcomes meaning there was imprecision and uncertainty in these results.
- 28 Uncertainty in the direction of the effect was also observed for quality of life outcomes. Given
- the limited evidence and the variation in current clinical practice, the committee could not make a recommendation for these agents and instead made a research recommendation
- 31 that included ACE inhibitors, beta-blockers and diuretics in adults with severe aortic stenosis
- 32 (see Appendix J.1.5 for details).

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## 2.1.6.3.2 Aortic regurgitation

- 34 There was insufficient evidence to draw conclusions about the relative benefits and harms of
- 35 ACE-I, ARB, beta-blockers and calcium channel blockers based on the evidence available.
- 36 The results indicated that there may be a clinically important benefit of ACE-inhibitors and
- 37 calcium channel blockers on the progression of heart valve disease on imaging, and a
- 38 benefit of calcium channel blockers for onset of symptoms and need for heart valve
- intervention. However, the evidence was graded low to very low quality for all outcomes and
- 40 comparisons and was based on a small number of studies with a small number of
- 41 participants, which the committee agreed gave imprecise estimates and was insufficient to
- show a true benefit or harm. They noted that a lot of these studies are historical and so may
- 43 not reflect current practice. Particularly they noted this for the comparison of digoxin to
- 44 calcium channel blockers, where the dose of digoxin was a higher dose than is used in
- modern practice, and may influence the results in this group. They further noted that digoxin
- 46 is not used currently for a ortic regurgitation. The presence of one study using it in this
- 47 population was explained by the fact that it is an old study and in the past digoxin was seen
- 48 as a possible treatment for many heart conditions but this is no longer the case in aortic
- 49 regurgitation. The committee also highlighted the lack of any placebo-controlled trials for
- 50 ARBs and digoxin. Instead of recommending any treatment (and due to variation in current
- 51 clinical practice), the committee made a research recommendation for more evidence, which
- 52 included ACE inhibitors, ARBs, beta-blockers and calcium channel blockers in adults with

- 1 aortic regurgitation (see Appendix J.1.1 for details). Digoxin was not included in this research
- 2 recommendation because it is no longer used in aortic regurgitation.

# 3 **2.1.6.3.3 Primary mitral regurgitation**

- 4 There was insufficient evidence to draw conclusions about the relative benefits and harms of
- 5 ACE-I and beta-blockers. The results suggested that ACE inhibitors may be beneficial for
- 6 preventing symptom onset, with a potential harm from adverse events. However, this was
- 7 based on two studies, with evidence being graded very low quality and the studies having
- 8 very small populations that did not report enough events to determine clinical importance,
- 9 resulting in very serious imprecision in the estimate of effect. The committee noted that this
- 10 population was younger than that which would be seen on average in the UK, and so may
- 11 not be representative. Instead of recommending any treatment (and due to variation in
- 12 current clinical practice), the committee made a research recommendation for more
- evidence, which included ACE inhibitors, beta-blockers and diuretics in adults with primary
- 14 severe mitral regurgitation (see Appendix J.1.10 for details).

# 15 **2.1.6.3.4 Key uncertainties**

- 16 There was no clinical evidence for mitral stenosis and tricuspid regurgitation. The committee
- 17 agreed that current practice does involve the use of pharmacological agents in tricuspid
- regurgitation (for example, diuretics) and were disappointed at the absence of evidence.
- 19 There was a lack of consensus about what treatment was appropriate and variation in
- 20 practice, meaning consensus recommendations could not be made for these populations.
- 21 Research recommendations were also not made for these populations as areas within
- 22 pharmacological treatment that were considered to be most feasible and useful were
- 23 prioritised for research recommendations.
- 24 There was insufficient evidence to make recommendations for the majority of these
- conditions. This is a key area of concern in current UK practice as there is uncertainty about
- 26 whether pharmacological management is required for people with heart valve disease to
- 27 prevent progression or delay consequences of the disease and the effect of pharmacological
- 28 treatment given for other conditions in those that also have heart valve disease. More
- specifically, there is uncertainty as to whether medications for the management of systemic
- 30 hypertensions are more poorly tolerated in the presence of valve disease. In addition, there
- 31 is uncertainty as to whether medications will delay the consequences of valve disease, for
- 32 example symptoms. Research recommendations were made covering some areas where
- 33 recommendations could not be made, however these were prioritised to the areas (aortic
- 34 regurgitation, severe aortic stenosis and severe primary mitral regurgitation) thought to be
- 35 most feasible and useful.

# 2.1.7 Cost effectiveness and resource use

- 38 No economic evaluations were found for this review question. The unit costs for the relevant
- 39 drug classes used to the treat heart failure without concomitant heart valve disease were
- 40 presented.
- 41 A cross-reference was made to the NICE guideline on lipid modification.
- 42 Due to a lack of clinical and economic evidence research recommendations were made for
- 43 all the other medicines.

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# 1 2.1.8 Other factors the committee took into account

- 2 Based on the recommendation, the committee agreed that there would not be a significant
- 3 effect on current practice. Based on the economic evidence, there would not be a substantial
- 4 cost implication from the use of statins.

5

# 6 2.1.9 Recommendations supported by this evidence review

- 7 This evidence review supports recommendation 1.2.1 and the research recommendations on
- 8 pharmacological management.

9

10

# 3.1 Review question: In adults with heart failure and

- 2 concomitant heart valve disease, what is the clinical and
- 3 cost effectiveness of ACE inhibitors, ARBs, beta blockers,
- 4 calcium channel blockers, digoxin, diuretics and nitrates to
- 5 improve clinical outcome?

# 3.1.d PICO table

7 For full details see the review protocol in Appendix A:.

# 8 Table 17: PICO characteristics of review question

Table 17: PICO c	haracteristics of review question
Population	Adults aged 18 years and over with diagnosed heart failure and heart valve disease of at least moderate severity stratified by type:  Primary aortic [including bicuspid] stenosis  Primary aortic regurgitation  Primary mitral stenosis  Primary mitral regurgitation  Primary tricuspid regurgitation  Secondary heart valve disease – mitral regurgitation or tricuspid regurgitation
Interventions	<ul> <li>Angiotensin-converting enzyme (ACE) inhibitors</li> <li>Angiotensin-II receptor blockers (ARBs)</li> <li>Beta blockers</li> <li>Calcium channel blockers (excluded for aortic stenosis)</li> <li>Digoxin</li> <li>Diuretics</li> <li>Nitrates (including nitroprusside)</li> <li>Any combination of 2 or more of the above</li> </ul>
Comparisons	<ul> <li>Placebo or no treatment</li> <li>Usual care (e.g. following standard heart failure guidelines: ACE + beta-blocker + diuretic)</li> <li>Other active comparator listed above, including combinations</li> </ul>
Outcomes	<ul> <li>All-cause mortality at 12 months (dichotomous)</li> <li>Cardiac mortality at 12 months (dichotomous)</li> <li>Hospital admission due to heart failure at 12 months (dichotomous)</li> <li>Health-related quality of life at 6 months and 12 months (continuous)</li> <li>Exercise tolerance reported as any of the following (in order of relevance): <ul> <li>Treadmill exercise time (duration)</li> <li>Time to near maximal dyspnoea</li> <li>6-minute walk test</li> <li>Borg dyspnoea index (continuous, final values or change scores)</li> </ul> </li> <li>Need for heart valve intervention (surgical or transcatheter) within 12 months (dichotomous)</li> <li>Withdrawal from the trial due to adverse events at 6 months and 12 months (dichotomous)</li> </ul>
Study design	Randomised control trials (RCTs) or systematic reviews of RCTs, including crossover trials

## 3.1.2 Clinical evidence

### 3.1.22 Included studies

- Ten studies were included in the review; 6, 15, 29, 38, 67, 87, 90, 123, 147, 150 these are summarised in 3
- Table 18 below. Evidence from these studies is summarised in the clinical evidence 4
- 5 summary Tables 19 to 24 below.
- 6 Evidence was only available for the following comparisons:
- 7 · Primary aortic stenosis:
- 8 o ACE-I versus placebo: 2 studies<sup>29, 38</sup>
- ARB versus placebo: 1 study<sup>67</sup> 9
- 10 Primary mitral stenosis
- Beta-blocker versus usual care: 2 studies<sup>87, 150</sup> 11
- o Beta-blocker versus placebo: 3 studies<sup>15, 90, 123</sup> 12
- Beta-blocker versus calcium channel blocker: 1 study<sup>6</sup> 13
- Secondary heart valve disease (mitral regurgitation or tricuspid regurgitation) 14
- ACE-I versus placebo: 1 study<sup>147</sup> 15
- No relevant RCTs for primary aortic, mitral or tricuspid regurgitation were identified. 17
- See also the study selection flow chart in Appendix C:, study evidence tables in Appendix D:, 18
- forest plots in Appendix E: and GRADE tables in Appendix F:. 19
- 20 Some of the studies included populations that did not directly match our protocol, as follows:
- Age Klein 1985<sup>87</sup> and Kumar 1994<sup>90</sup> included participants under the age of 18 21
- Severity of heart valve disease Klein 1985<sup>87</sup>, Patel 1995<sup>123</sup> and Seneviratne 1994<sup>147</sup> did 22 not provide information on or to determine the severity of heart valve disease for their 23
- 24 population.

#### 3.1.**2**.52 **Excluded studies**

26 See the excluded studies list in Appendix I:.

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# 3.1.2.8 Summary of clinical studies included in the evidence review

# Table 18: Summary of studies included in the evidence review

Study	Intervention and comparison	Population	Outcomes	Comments
Alan 2002 <sup>6</sup>	Beta blocker (metoprolol) Initially 5mg intravenous, followed by 50mg orally twice daily  Calcium channel blocker (diltiazem) Initially 25mg intravenously, followed by 60mg orally three times daily	Adults with symptomatic mild-to-moderate mitral stenosis receiving chronic maintenance therapy.  New York Heart Association (NYHA) class II and III.  Mean ± SD age 38±6.8.  Turkey	Exercise tolerance at 3 months Withdrawal due to adverse events at 3 months	Proportion with mild mitral stenosis not stated. Baseline total effort time not matched
Bassan 1987 <sup>15</sup>	Beta blocker (propranolol) 40mg orally twice or three times daily (dependent on weight).  Placebo	Adults with moderate symptomatic isolated mitral stenosis waiting for surgery (5 had surgery 1-24 months post-intervention).  Median mitral valve area 1.1cm² (severity of valve disease not stated directly).  NYHA class II and III.  Mean age 38.7 (range: 19-56)	Exercise tolerance at 1 week	Crossover RCT – insufficient data were available to account for the within-patient correlation and so the data were analysed as if it were a parallel trial
Chockalingam 2004 <sup>29</sup>	ACE inhibitor (enalapril) 2.5mg twice daily titrated up to 10mg twice daily over 2 weeks  Placebo	Adults with symptomatic severe aortic stenosis waiting for surgery or unwilling to have surgery.  NYHA class III and IV.	Exercise tolerance at 4 weeks Withdrawal due to adverse events at 3 months	

Study	Intervention and comparison	Population	Outcomes	Comments
		Mean ± SD age Intervention group: 43±11, Control group: 46±12		
Dalsgaard 2014 <sup>38</sup>	ACE inhibitor (trandolapril) Daily increasing doses up to the maximum tolerated dose (maximum: 2mg).  Placebo	Adults with severe symptomatic (32) and asymptomatic (12) aortic stenosis waiting for surgery. NYHA classes II to IV.  Mean ± SD age 69.9±8.3.  Denmark	Exercise tolerance at 3 days Withdrawal due to adverse events at 8 weeks	30 patients had comorbidities (including hypertension, ischaemic heart disease, and diabetes mellitus).
Helske-Suihko 2015 <sup>67</sup>	ARB (candesartan) 8mg once daily for 2 weeks, then 16mg once daily until 3 days before they have valve surgery (mean: 5.4 months).  Placebo	Adults with symptomatic severe aortic stenosis waiting for surgery.  Majority NYHA class II.  Mean ± SD age Intervention: 73±9, control: 70±12.  10% were in atrial fibrillation or pacemaker rhythm  Finland	Hospitalisation due to heart failure at 2-12 months (mean: 5.4 months)  Exercise tolerance at 2-12 months (mean: 5.4 months)  Withdrawal due to adverse events at 2-12 months (mean: 5.4 months)  All-cause mortality at 2-12 months (mean: 5.4 months)	Reports the majority of patients had symptoms equivalent to NYHA class II. However, then selectively reports the proportion in class I/II vs. class III.  All analysed participants underwent valve replacement as part of the study protocol (so this is not reported as an outcome)
Klein 1985 <sup>87</sup>	Beta blockers (atenolol) 100mg once daily for 2 weeks.  Placebo	People (age range 15-35 years) with symptomatic significant isolated mitral stenosis in sinus rhythm receiving chronic maintenance therapy.  Severity of valve disease not stated.	Exercise tolerance at 2 weeks	Crossover RCT – insufficient data were available to account for the within-patient correlation and so the data were analysed as if it were a parallel trial.  Unclear what proportion <18 years of age were included.

Study	Intervention and comparison	Population	Outcomes	Comments
		NYHA class II and III. Age range 15-35 years. South Africa		
Kumar 1994 <sup>90</sup>			Exercise tolerance at 6 months	Unclear if any <18 years of age were included.
Patel 1995 <sup>123</sup>	Beta blockers (acebutolol or atenolol) Acebutolol 400mg once daily or atenolol 100mg daily for 1 week.  Placebo	People with symptomatic isolated mitral stenosis admitted for percutaneous mitral valvotomy.  Severity of valve disease not stated.  NYHA class II and III.  Mean age 28(range 17-51 years).	Exercise tolerance at 4 months	Crossover RCT – insufficient data were available to account for the within-patient correlation and so the data were analysed as if it were a parallel trial Unclear what proportion <18 years of age were included.

Study	Intervention and comparison	Population	Outcomes	Comments
		South Africa		
Seneviratne 1994 <sup>147</sup>	ACE inhibitors (captopril) 6.25mg twice daily, increasing to 12.5mg twice daily after 4 weeks, increasing to 25mg twice daily after 8 weeks, increasing to 50mg twice daily at 12 weeks.  Placebo	Adults with symptomatic secondary mitral regurgitation receiving chronic maintenance therapy.  Severity of mitral regurgitation was not stated.  NYHA class II and III.  Mean ± SD age Captopril mean: 72.3±5.4, Placebo mean: 71.5±7.2).	Quality of life at 12 weeks Cardiac mortality at 12 weeks Withdrawal due to adverse events at 12 weeks	Study funded by industry
Shu 2005 <sup>150</sup>	Beta blockers (bisoprolol) Initial dose 1.25mg/day. Recommended maximum dose 10mg/day. Gradual titration over 3-5 days by 2-3 weeks.  Usual care All patients received warfarin and basic therapy with one of the following: a diuretic, digoxin, ACE-inhibitors (or ARBs if contraindicated), or nitrates.	Adults with symptomatic significant mitral stenosis or aortic lesions and mitral regurgitation from uncorrected rheumatic heart valvular disease and atrial fibrillation receiving chronic maintenance therapy.  Significant valve disease defined as: aortic stenosis with a gradient greater than 20mmHg; mitral stenosis with a valve area of less than 1.5cm², or mitral valve regurgitation lesions of at least moderate severity. Approximately 50% had significant mitral stenosis.	Hospitalisation due to heart failure at 12 months Exercise tolerance at 6-12 months Withdrawal due to adverse events at 12 months	

Study	Intervention and comparison	Population	Outcomes	Comments
		Mean age Intervention: 40.6±6.8; Control: 43.5±7.4).		
		China		

1 See Appendix D: for full evidence tables.

# 3.1.22 Quality assessment of clinical studies included in the evidence review

# 3 1.2.431 Primary aortic stenosis

Table 19: Clinical evidence summary: ACE-I versus placebo in primary aortic stenosis

	No of			Anticipated absolute effects		
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with Placebo	Risk difference with ACE-I (95% CI)	
Exercise tolerance: change in exercise duration (minutes) semi-supine cycle exercise test	43 (1 study) 3 days	⊕⊕⊝ LOW <sub>1,2.3</sub> due to risk of bias, indirectness		The mean change in exercise duration (minutes) in the control groups was 0.2 minutes	The mean change in exercise duration (minutes) in the intervention groups was 0 higher (0.31 lower to 0.31 higher)	
Exercise tolerance: 6- minute walk distance (meters)	52 (1 study) 4 weeks	⊕⊕⊖⊖ LOW <sub>1,4,5</sub> due to risk of bias, imprecision		The mean exercise tolerance: 6-minute walk distance (meters) in the control groups was 376 meters	The mean exercise tolerance: 6-minute walk distance (meters) in the intervention groups was 26 higher (68.89 lower to 120.89 higher)	
Withdrawal due to adverse events	100 (2 studies) 2-3 months	⊕⊖⊖ VERY LOW <sub>6,7,8</sub> due to risk of bias, indirectness, imprecision	Peto OR 2.18 (0.34 to 14.17)	26 per 1000	29 more per 1000 (from 17 fewer to 248 more)	

<sup>1</sup> Downgraded by 1 increment because the evidence was at high risk of bias

<sup>2</sup> Downgraded by 1 increment because the mean follow-up period was less than 1 month

	No of			Anticipated absolute effects	
	Participants (studies)	Quality of the evidence	Relative effect		
Outcomes	Follow up	(GRADE)	(95% CI)	Risk with Placebo	Risk difference with ACE-I (95% CI)

- 3 MIDs used to assess imprecision were ±1.0
- 4 Downgraded by 1 increment because the confidence interval crossed one MID
- 4 Downgraded by 1 increment because the majority of evidence was at high risk of bias
- 5 MIDs used to assess imprecision were ±76.0
- 6 Downgraded by 1 increment because the mean follow-up period was less than 3 months
- 7 Downgraded by 2 increments because the confidence interval crossed both MIDs
- 8 Downgraded by 1 increment because the majority of evidence was at high risk of bias

# 1 Table 20: Clinical evidence summary: ARB versus placebo

	No of Participants (studies)	Quality of the evidence	Relative effect	Anticipated absolute effects	
Outcomes	Follow up	(GRADE)	(95% CI)	Risk with Placebo	Risk difference with ARB (95% CI)
All-cause mortality	51 (1 study) 2-12 months	⊕⊖⊖ VERY LOW <sub>1,2</sub> due to risk of bias, imprecision	Peto OR 0.14 (0 to 7.09)	39 per 1000	39 fewer per 1000 (from 140 fewer to 63 more) <sub>7</sub>
Acute heart failure	51 (1 study) 2-12 months	⊕⊖⊖ VERY LOW <sub>1,2,3</sub> due to risk of bias, indirectness, imprecision	Peto OR 7.69 (0.15 to 387.87)	0 per 1000	40 more per 1000 (from 63 fewer to 143 more) <sub>7</sub>
Exercise tolerance: change from baseline 6-minute walking distance (meters)	43 (1 study) 2-12 months	⊕⊕⊕⊝ MODERATE <sub>4,5</sub> due to risk of bias		The mean exercise tolerance: change from baseline 6-minute walking	The mean change from baseline in 6-minute walking distance in the intervention groups was

Outcomes	No of Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Anticipated absolute effects  Risk with Placebo Risk difference with ARB (95% CI)		
				distance in the control groups was -2 meters	18 metres lower (48.74 lower to 12.74 higher)	
Withdrawal due to adverse events	51 (1 study) 2-12 months	⊕⊖⊖⊖ VERY LOW <sub>1,6</sub> due to risk of bias, imprecision	RR 1.04 (0.16 to 6.83)	77 per 1000	3 more per 1000 (from 65 fewer to 449 more)	

- 1 Downgraded by 2 increments because the evidence was at very high risk of bias
- 2 Downgraded by 1 increment because the confidence interval crossed one MID
- 3 Downgraded by 1 increment because of uncertainty as to the aetiology of reported acute heart failure
- 4 Downgraded by 1 increment because the evidence was at high risk of bias
- 5 MIDs used to assess imprecision were ±74.0
- 6 Downgraded by 2 increments because the confidence interval crossed both MIDs
- 7 Absolute effect calculated manually using risk difference as zero events in one arm of the study

# 3₹1.2.412 Primary mitral stenosis

Table 21: Clinical evidence summary: Beta-blocker versus usual care

	No of Participants Quality of the			Anticipated absolute effects		
Outcomes	(studies) Follow up	evidence (GRADE)	Relative effect (95% CI)	Risk with Usual care	Risk difference with Beta-blocker (95% CI)	
Hospitalisation due to heart failure	67 (1 study) 12 months	⊕⊖⊖ VERY LOW <sub>1,2</sub> due to risk of bias, imprecision	RR 0.31 (0.09 to 1.02)	294 per 1000	203 fewer per 1000 (from 268 fewer to 6 more)	
Exercise tolerance: 6- minute walking distance (meters)	67 (1 study) 6-12 months	⊕⊕⊖⊖ LOW <sub>1,3</sub> due to risk of bias		The mean exercise tolerance: 6-minute walking distance in the	The mean exercise tolerance: 6-minute walking distance in the intervention groups was	

Outcomes	No of Participants	Quality of the evidence (GRADE)		Anticipated absolute effects		
	(studies) Follow up		Relative effect (95% CI)	Risk with Usual care	Risk difference with Beta-blocker (95% CI)	
				control groups was 290 meters	133 meters higher (121.49 to 144.51 higher)	
Withdrawal due to adverse events (weakness, dizziness, dyspnoea)	88 (1 study) 12 months	⊕⊕⊝⊝ LOW₁ due to risk of bias	Peto OR 8.14 (1.35 to 48.97)	0 per 1000	114 more per 1000 (from 13 more to 214 more) <sub>4</sub>	

- 1 Downgraded by 2 increments because the evidence was at very high risk of bias 2 Downgraded by 1 increment because the confidence interval crossed one MID
- 3 MIDs used to assess imprecision were ±15.0
- 4 Absolute effect calculated manually as zero events in one arm of the study

# Table 22: Clinical evidence summary: Beta-blocker versus placebo

No of Participants	Participants	Quality of the evidence (GRADE)	Relative effect (95% CI)	Anticipated absolute effects		
Outcomes	(studies) Follow up			Risk with Placebo	Risk difference with Beta blocker (95% CI)	
Exercise tolerance: treadmill exercise time (minutes) to exhaustion	84 (3 studies) 1-4 weeks	⊕⊖⊖ VERY LOW <sub>1,2,3,4,5</sub> due to risk of bias, inconsistency, indirectness, imprecision,		The mean treadmill exercise time (minutes) to exhaustion in the control groups was 8.1 minutes	The mean treadmill exercise time (minutes) to exhaustion in the intervention groups was 0.33 higher (1.09 lower to 1.75 higher)	
Exercise tolerance: Pulmonary capillary	26 (1 study) 6 months	⊕⊕⊖⊖ LOW <sub>6,7,8</sub>		The mean pulmonary capillary wedge pressure after exercise in the control group was	The mean pulmonary capillary wedge pressure after exercise in the intervention groups was	

	No of Participants	Quality of the evidence (GRADE)	Relative effect (95% CI)	Anticipated absolute effects		
Outcomes	(studies) Follow up			Risk with Placebo	Risk difference with Beta blocker (95% CI)	
wedge pressure after exercise		due to risk of bias, indirectness		50.5	14.8 lower (21.71 to 7.89 lower)	

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- 1 Downgraded by 2 increments because the majority of the evidence was at very high risk of bias
- <sub>2</sub> Downgraded by one increment because the I2 = 74% and heterogeneity was not explained by subgroup analyses.
- 3 Downgraded by 1 increment because the mean follow-up period is less than 1 month
- 4 Downgraded by 1 increment because the confidence interval crossed one MID
- 5 MIDs used to assess imprecision were ±0.9
- 6 Downgraded by 1 increment because the evidence was at high risk of bias
- 7 Downgraded by 1 increment because the outcome is a surrogate measure
- 8 MIDs used to assess imprecision were ±5.35

# Table 23: Clinical evidence summary: Beta-blocker versus calcium channel blocker

	No of Participants	Quality of	Relative effect (95% CI)	Anticipated absolute effects	
Outcomes	(studies) Follow up	the evidence (GRADE)		Risk with calcium channel blocker	Risk difference with beta blocker (95% CI)
Exercise tolerance: total effort time on treadmill exercise test	80 (1 study) 3 months	⊕⊖⊖ VERY LOW <sub>1,2,3</sub> due to risk of bias, imprecision		The mean exercise tolerance: total effort time on treadmill exercise test in the control groups was 570 seconds <sub>7</sub>	The mean total effort time on treadmill exercise test in the intervention groups was 50 seconds lower (97.99 to 2.01 lower)
Withdrawal due to adverse events	80 (1 study) 3 months	⊕⊕⊖⊖ LOW <sub>4,6</sub> due to risk of bias, imprecision	RD 0 (- 0.048 to 0.048)	0 per 1000	0 fewer per 1000 (from 48 fewer to 48 more)5

<sup>1</sup> Downgraded by 2 increments because the evidence was at very high risk of bias: baseline total effort time not matched – beta-blocker: 452±120; calcium-channel blocker: 534±120

<sup>2</sup> Downgraded by 1 increment because the confidence interval crossed one MID

	No of Participants	Quality of	Relative	Anticipated absolute effects		
Outcomes	(studies) the e	the evidence (GRADE)	effect (95% CI)	Risk with calcium channel blocker	Risk difference with beta blocker (95% CI)	

- 3 MIDs used to assess imprecision were ±60.0
- 4 Downgraded by 1 increment because the evidence was at high risk of bias
- 5 Absolute effect calculated manually as zero events in both arms of the study
- 6 Downgraded by 1 increment because sample size was >70 and <350 (imprecision was assessed based on sample size as zero events in both arms of the study)
- 7 Baseline total effort time not matched

# 3 1.2.413 Secondary heart valve disease (mitral regurgitation or tricuspid regurgitation)

Table 24: Clinical evidence summary: ACE-I versus placebo in secondary heart valve disease

		Relative effect	Anticipated absolute effects		
Outcomes	Follow up	(GRADE)	(95% CI)	Risk with Placebo	Risk difference with ACE-I (95% CI)
Cardiac mortality	28 (1 study) 12 weeks	⊕⊕⊝⊝ LOW <sub>1,2</sub> due to risk of bias, imprecision	OR 0.14 (0 to 6.82)	71 per 1000	71 fewer per 1000 (from 248 fewer to 106 more)6
Quality of life: Duke activity index score Scale: 2.75 to 58.2 (high is good outcome)	23 (1 study) 12 weeks	⊕⊖⊖ VERY LOW <sub>2,3,4,5</sub> due to risk of bias, indirectness, imprecision		The mean Duke activity index score in the control group was 22.3	The mean Duke activity index score in the intervention groups was 6.7 higher (0.97 lower to 14.37 higher)
Withdrawal due to adverse events	27 (1 study) 3 months	⊕⊖⊖ VERY LOW <sub>1,8</sub> due to risk of bias, imprecision	RD 0 (- 0.133 to 0.133)	0 per 1000	0 fewer per 1000 (from 133 fewer to 133 more)7

<sup>1</sup> Downgraded by 1 increment because the evidence was at high risk of bias

<sup>2</sup> Downgraded by 1 increment because the confidence interval crossed one MID

	(studies) evi	Quality of the evidence (GRADE)	Relative effect (95% CI)	Anticipated absolute effects		
Outcomes				Risk with Placebo	Risk difference with ACE-I (95% CI)	

- 3 Downgraded by 2 increments because the evidence was at very high risk of bias
- 4 Downgraded by 1 increment because the reported measure only reports physical activity rather than other aspects of quality of life
- 5 MIDs used to assess imprecision were ±4.7
- 6 Absolute effect calculated manually using risk difference as zero events in one arm of the study 7 Absolute effect calculated manually as zero events in both arms of the study
- 8 Downgraded by 2 increments because sample size was <70 (imprecision was assessed based on sample size as zero events in both arms of the study)
- See Appendix F: for full GRADE tables.
- 2
- 3

# 3.1.3 Economic evidence

## 3.1.32 Included studies

3 No health economic studies were included.

# 3.1.3.2 Excluded studies

- 5 No relevant health economic studies were excluded due to assessment of limited
- 6 applicability or methodological limitations.
- 7 See also the health economic study selection flow chart in Appendix G:.

8

# 3.1.3.8 Summary of studies included in the economic evidence review

2

No economic studies were included in this review.

4

# 3.1.3.4 Unit costs

4

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The following relevant unit costs have been included to inform the committee of the cost implications of different pharmacological management strategies.

Table 25. Unit costs for different drugs used for pharmacological management of people with heart failure and concomitant heart valve disease

ACE inhibitors ramipril 1.25mg £ 2.5mg £	0.07 0.15 0.17
	0.17
5mg £	
10mg £	0.18
captopril 12.5mg £	0.02
25mg £	0.01
50mg £	0.03
enalapril 2.5mg £	0.18
5mg, 10mg, 20mg £	0.06
lisinopril 2.5mg, 5mg, 10mg, 20mg £	0.03
quinapril 2.5mg, 5mg, 10mg £	0.31
20mg £	0.39
40mg £	0.13
fosinopril 10mg £	0.15
20mg £	0.14
Angiotensin II candesartan cilexitil 2mg £	0.22
receptor blockers (ARBs) 4mg	0.08
£ 8mg	0.04
16mg £	0.06
32mg £	0.06
losartan 12.5mg £	0.11
25mg £	0.12
50mg £	0.07
100mg £	0.07
Beta blockers bisoprolol 1.25mg £	0.03
3.75mg £	0.03
5mg £	0.02
10mg £	0.03
carvedilol 3.125mg £	0.03
6.25mg £	0.03
12.5mg £	0.03
25mg £	0.04
nebivolol 2.5mg £	0.42

		5mg	£	0.16
		10mg	£	0.92
Diuretics	furosemide	20mg tablet	£	0.05
		40mg tablet	£	0.07
		10 mg per 1 ml solution for injection	£	1.74
	bumetanide	1mg tablet	£	0.05
		5mg tablet	£	0.25
	torasemide	2.5mg tablet	£	0.14
		5mg	£	0.20
		10mg	£	0.29
Calcium channel blockers	amlopodine	5mg, 10mg	£	0.03
Digoxin	-	62.5 micrograms	£	0.05
		125 micrograms	£	0.05
Nitrates	Isosorbide dinitrate	10mg	£	0.24

1 Source: BNF 2018<sup>79</sup>

## 3.1.24 Evidence statements

# 3.1.43 Clinical evidence statements

4 See the summary of evidence in Tables 19-24.

## 3.1.4.2 Health economic evidence statements

6 7

No relevant economic evaluations were identified.

8

# 3.1.5 The committee's discussion of the evidence

# 3.11.6 Interpreting the evidence

# 3.1.6.1 The outcomes that matter most

- 12 The critical outcomes were all-cause mortality, cardiac mortality, hospital admission due to
- heart failure and health-related quality of life. Important outcomes were exercise tolerance,
- 14 need for heart valve intervention (surgical or transcatheter) and withdrawal from the study
- 15 due to adverse events.
- 16 Physiological outcomes were not included in the protocol as they are not clinically relevant
- endpoints, and the outcomes they aim to predict that are important to patients are captured
- 18 by the included outcomes. However, one study reported pulmonary wedge pressure
- 19 following exercise for the comparison of beta-blockers compared to placebo in primary mitral
- 20 stenosis, which was included but downgraded for indirectness as it is a surrogate measure of
- 21 exercise tolerance.
- 22 There was very limited evidence, especially for health-related quality of life. The need for
- 23 heart valve intervention was not reported in any of the studies.

# 3.1.6.2 The quality of the evidence

- 2 No relevant RCTs for primary aortic, mitral or tricuspid regurgitation were identified. Ten
- 3 RCTs were included in this review and evidence was only available for the following
- 4 comparisons:

9

- Primary aortic stenosis:
- 6 o ACE-I versus placebo
- 7 o ARB versus placebo
- Primary mitral stenosis
  - Beta-blocker versus usual care
- o Beta-blocker versus placebo
- 11 o Beta-blocker versus calcium channel blocker
- Secondary heart valve disease (mitral regurgitation or tricuspid regurgitation)
- o ACE-I versus placebo
- 14 Evidence ranged from moderate to very low quality, with the majority of the evidence being of
- low or very low quality. Evidence was mainly downgraded due to risk of bias and imprecision.
- Analyses frequently included only a small number of participants and had low event rates
- 17 resulting in great uncertainty. Additionally, some evidence was considered to be indirect
- because of the inclusion of populations that did not reflect those seen in UK practice or
- reporting at time points shorter than 3 months (or 1 month for exercise tolerance).

## 3.1.63 Benefits and harms

# 21 Primary aortic stenosis

- 22 There was insufficient evidence to draw conclusions about the relative benefits and harms of
- ACE-I or ARB compared with placebo, as there was only one very small study identified for
- 24 each outcome and comparison, and no other comparisons were available for this stratum. No
- 25 clinically important differences were seen for any of the reported outcomes, though
- 26 uncertainty was observed for all outcomes and the majority of the evidence for all
- 27 comparisons was graded low to very low quality. Due to variation in current clinical practice
- 28 the committee were unable to make consensus recommendations. A research
- 29 recommendation was therefore made to investigate the clinical and cost-effectiveness of
- 30 pharmacological management of heart failure in adults with severe aortic stenosis (see
- 31 Appendix J.2.6 for details). This research recommendation was also applied to the severe
- 32 aortic regurgitation and severe mitral regurgitation populations, as no evidence was identified
- 33 for these populations.

# 34 Primary mitral stenosis

- 35 For the comparison of beta blocker vs usual care (warfarin and basic therapy with either a
- diuretic, digoxin, ACE-inhibitors (or ARBs if contraindicated), or nitrates) clinically important
- 37 benefits of beta blockers were seen for reduced hospitalisation due to heart failure and
- increased exercise tolerance, though some uncertainty in the direction of the effect was
- 39 observed for the hospitalisation due to heart failure outcome. However, there was also a
- 40 clinically significant increase in the rate of withdrawals due to adverse events, namely
- 41 weakness, dizziness and dyspnoea. This was based on evidence from a single, small study
- 42 with evidence graded low to very low quality. Beta-blockers are widely used in patients with
- 43 heart failure due to reduced ventricular systolic function and in patients with coronary artery
- 44 disease and the same adverse events do not represent sufficient reason to negate potential
- 45 benefit. The committee noted that in the study all participants were also in atrial fibrillation.
- 46 The evidence for the comparisons of beta blockers with placebo and calcium channel
- 47 blockers was limited, based on factors such as inconsistency in results between studies and

1 quality of the evidence being graded low to very low quality with very small population sizes, 2 and it was not possible to draw any conclusions from the data. For the comparison between 3 beta-blockers and calcium channel blockers, there was some suggestion of worse exercise 4 tolerance with beta-blockers, but this evidence was considered to be limited given it was 5 based on a single, small study and the size of the effect was uncertain based on confidence 6 intervals, with the quality also being graded very low. For the comparison between beta-7 blockers and placebo, it was noted that the heterogeneity for the outcome of exercise 8 tolerance could have been due to heart valve disease severity, as the study showing a 9 benefit of beta blockers was in a population with 'significant' mitral stenosis while the other 10 studies included moderate severity or did not specify the severity. However, due to the limited number of studies and the poorly defined populations it was not possible to assess 11 12 this formally. It was also noted that the study showing a benefit included only people in sinus 13 rhythm, while the others did not report the numbers in sinus rhythm or atrial fibrillation. This 14 highlights the need to address whether beta blockers are effective in both sinus rhythm and 15 atrial fibrillation, and a research recommendation was made to encourage research in this 16 area (see Appendix J.2.1 for details).

17 The committee noted that all studies in this stratum related to a much younger population 18 than the cohort seen in UK clinical practice and was often due to rheumatic fever, excluding 19 those aged over 75 years. This is distinct from the cases seen in the UK which are commonly 20 in older adults with calcific mitral stenosis. For this reason, the research recommendation 21 described above on beta-blockers in mitral stenosis for those in sinus rhythm and those in 22 atrial fibrillation was limited to adults ≥75 years in order to provide some direct evidence in 23 this age group. The committee also acknowledged that one key study only included people in 24 atrial fibrillation. However, the committee agreed that the findings align with their clinical 25 experience in the UK population and can be extrapolated to this group. It is also plausible 26 that lowering the heart rate in mitral stenosis should produce a benefit for patients.

27 Therefore, based on the evidence, supported by their clinical experience, the committee 28 made a recommendation to consider beta blockers in people with moderate to severe mitral 29 stenosis and concomitant heart failure. The recommendation was a consider 30 recommendation based on the limitations associated with the included evidence, including 31 the small study size, quality of the evidence being graded low to very low and uncertainty in 32 the direction of the effect for many some outcomes. A separate recommendation was made 33 rather than referring to the NICE chronic heart failure guideline as it was explained that in 34 cases where heart failure is due to the heart valve disease, reduced systolic function may not 35 be present so it would therefore not be appropriate to refer to this guideline. In addition, the research recommendation described in the previous paragraphs was also made to assess 36 37 the clinical and cost-effectiveness of beta-blockers in adults ≥75 years, both in sinus rhythm 38 and in atrial fibrillation, so that there is direct evidence for this older population with non-39 rheumatic/calcific mitral stenosis.

# Secondary mitral or tricuspid regurgitation

41 A single, very small trial was available comparing ACE-I vs placebo. There was a possible benefit for improved quality of life based on functional ability assessed in the Duke Activity 42 43 Index score, though there was uncertainty in the direction of the effect. However, overall there was insufficient evidence to draw conclusions, based on the size of the study, evidence 44 45 being graded low to very low quality and uncertainty in the effect estimates for the other 46 available outcomes of cardiac mortality and withdrawal due to adverse events. Due to 47 variation in current clinical practice the committee were unable to make consensus 48 recommendations.

## Key uncertainties

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Only 3 out of 6 of the studies in the mitral stenosis stratum specified whether the participants were in sinus rhythm or atrial fibrillation and none included older adults with calcific heart valve disease. Therefore, to inform future updates of this guidance the committee made a

- 1 research recommendation around beta blockers, the key pharmacological intervention in this
- 2 group, for older adults with non-rheumatic/calcific mitral stenosis including groups in sinus
- 3 rhythm and in atrial fibrillation. This is to encourage research to clarify whether this form of
- 4 pharmacological management is safe and effective in the population most relevant to UK
- 5 clinical practice, both in people in sinus rhythm and those in atrial fibrillation, as there is
- 6 currently no randomised evidence to answer these important clinical questions.
- 7 There was insufficient evidence to inform a recommendation for people with aortic stenosis.
- 8 This is a key area of concern in current UK practice as there is uncertainty about whether
- 9 pharmacological management in severe aortic stenosis is appropriate. Therefore, a research
- 10 recommendation was made to encourage research into the clinical and cost effectiveness of
- 11 pharmacological management of heart failure in adults with severe aortic stenosis. This
- 12 research recommendation also applied to those with severe aortic regurgitation and severe
- mitral regurgitation, as no evidence was identified for these populations.
- 14 Although there was an absence of evidence for other areas included in the review protocol,
- including primary tricuspid regurgitation and secondary mitral and tricuspid regurgitation,
- 16 consensus recommendations could not be made due to variation in practice and research
- 17 recommendations were prioritised to the areas thought to be most feasible and useful.

# 3.118 Cost effectiveness and resource use

- 19 No economic evaluations were found for this review question. The unit costs for the relevant
- 20 drug classes used to treat heart failure with concomitant heart valve disease were presented.
- 21 The committee agreed that due to the low cost of all relevant drugs, the interventions costs
- 22 for pharmacological management were unlikely to differ substantially from one drug class to
- 23 another.
- 24 The clinical review demonstrated that using beta blockers to manage mitral stenosis may be
- 25 associated with reduced hospital readmissions (due to heart failure) compared to usual care
- 26 (warfarin and basic therapy with either a diuretic, digoxin, ACE-inhibitors (or ARBs if
- contraindicated), or nitrates), although there was uncertainty in the direction of effect for this
- outcome and evidence was only available from a single, small study with evidence graded
- 29 very low quality. Therefore, there may be cost savings by considering beta blockers for this
- 30 population over these comparators. However, some of these savings may be offset as
- 31 people will need to be monitored for adverse events and no studies reported this outcome for
- 32 beta-blockers compared with other comparators, such as placebo or calcium channel
- 33 blockers.

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- 34 Given that beta-blockers already form a large part of current practice for mitral stenosis, their
- recommendation is not likely to have a resource impact. There should be a focus on correct
- 36 titration of beta blockers as improper titration may increase the need for monitoring.

# 3.88 Other factors the committee took into account

- 39 The committee made a research recommendation on the pharmacological management in
- 40 adults with severe aortic stenosis, severe aortic regurgitation or severe mitral regurgitation to
- 41 address the lack of evidence in this area.
- When developing the protocol for this review the committee discussed the current focus on
- 43 the neprilysin inhibitor sacubitril in combination with valsartan in heart failure research. It was
- 44 noted that the current guideline must focus on the management of heart valve disease and
- 45 cannot assess all pharmacological management options for heart failure if not currently used
- in people with heart valve disease, as it was agreed that this combination is not commonly
- 47 used in heart valve disease. The committee were aware of the recommendations on the
- 48 pharmacological management of chronic heart failure in the NICE guideline on the chronic

### Heart valve disease: DRAFT FOR CONSULTATION 4 MIDs used to assess imprecision were ±4.0

- 1 heart failure (NG106). It was agreed that future updates of this guidance may be able to
- 2 assess the use of this drug combination in heart valve disease with concomitant heart failure.

#### 3.1.3 Recommendations supported by this evidence review

- 4 This evidence review supports recommendation 1.2.2 and the research recommendations on 5
  - pharmacological management.

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# **Appendices**

3

# 2 Appendix A: Review protocols

## A.4 Valve disease without heart failure

Table 26: Review protocol: pharmacological management of heart valve disease without concomitant heart failure

ID	Field	Content	
0.	PROSPERO registration number	Not registered	
1.	Review title	In adults with heart valve disease without concomitant heart failure, what is the clinical and cost effectiveness of alpha-blockers, angiotensin-converting enzyme (ACE) inhibitors, angiotensin-II receptor blockers (ARBs), beta blockers, calcium channel blockers, digoxin, diuretics, statins and nitrates to improve clinical outcome?	
2.	Review question	In adults with heart valve disease without concomitant heart failure, what is the clinical and cost effectiveness of alpha-blockers, ACE inhibitors, ARBs, beta blockers, calcium channel blockers, digoxin, diuretics, statins and nitrates to improve clinical outcome?	
3.	Objective	To assess the clinical and cost-effectiveness of pharmacological interventions individually and in combination to manage asymptomatic heart valve disease in adults.	
4.	Searches	The following databases will be searched:	
		Cochrane Central Register of Controlled Trials (CENTRAL)	
		Cochrane Database of Systematic Reviews (CDSR)	
		Embase	
		MEDLINE	

		Searches will be restricted by:
		English language studies
		Human studies
		Letters and comments are excluded
		Validated study filters for systematic reviews and RCTs
		No date restrictions applied
		The searches may be re-run 6 weeks before final committee meeting and further studies retrieved for inclusion if relevant.
		The full search strategies database will be published in the final review.
5.	Condition or domain being studied	Diagnosed heart valve disease in adults aged 18 years and over: Aortic (including bicuspid) stenosis, aortic regurgitation, mitral stenosis, mitral regurgitation and tricuspid regurgitation.
6.	Population	Inclusion:
		Adults aged 18 years and over with diagnosed heart valve disease of at least moderate severity stratified by type:
		Primary aortic [including bicuspid] stenosis
		Primary aortic regurgitation
		Primary mitral stenosis
		Primary mitral regurgitation
		Primary tricuspid regurgitation
		Secondary heart valve disease – mitral regurgitation or tricuspid regurgitation
		A study will be considered to cover a population with heart valve disease without concomitant heart failure if it meets all of the following criteria:
		Diagnosis of native heart valve disease

		<ul> <li>Asymptomatic or have only very mild/low-level symptoms that would not affect daily life (this would include those reported to be in class I of the NYHA classification)</li> <li>A normal LVEF</li> </ul> Include only first line use of pharmacological management options.
		Inclusion of indirect evidence from mixed populations, to be considered separately for each strata and intervention:
		Studies including adults with HVD where some also have concomitant heart failure will be included if <50% of the included patients had heart failure (those studies with ≥50% concomitant heart failure will be included in a separate review question focused on HVD and concomitant heart failure) .
7.	Intervention/Exposure/Test	Alpha blockers
		Angiotensin-converting enzyme (ACE) inhibitors
		Angiotensin-II receptor blockers (ARBs)
		Beta blockers
		Calcium channel blockers
		Digoxin
		• Diuretics
		Statins
		Nitrates (including nitroprusside)
		Any combination of 2 or more of the above
		Primary studies with a mixed intervention (some in the 'active' arm received the intervention of interest and some a different intervention) will be included if at least 90% received the intervention of interest.
		A class effect will be used for analysis, combining all interventions within each drug class (regardless of mode of delivery, and dose – as long as within the licensed range).
		For crossover studies, there is no lower limit for the washout period because drug effects do not persist once a person is no longer taking the drug.

8.	Comparator/Reference standard/Confounding factors	<ul> <li>Placebo or no treatment (usual care)</li> <li>Other active comparator listed above, including combinations</li> </ul>
9.	Types of study to be included	Randomised control trials (RCTs) or systematic reviews of RCTs, including crossover trials
		If no RCT data are available, observational data will not be considered for pharmacological interventions. This is due to the risk of confounding variables influencing the study results, reducing our confidence in the review results.
10.	Other exclusion criteria	Exclusion criteria:
		<ul> <li>Conference abstracts will be excluded because they are unlikely to contain enough information to assess whether the population matches the review question in terms of previous medication use, or enough detail on outcome definitions, or on the methodology to assess the risk of bias of the study.</li> </ul>
		Non-randomised studies / observational studies
		<ul> <li>Pharmacological management of HVD in adults with concomitant heart failure (where the proportion with heart failure is ≥50%)</li> </ul>
		Pharmacological management of mild HVD
		Pharmacological management in the intra- or post-operative period
		• Studies combining people with different types of valve disease (e.g. some with mitral regurgitation and some with aortic regurgitation).
		Pharmacological management in children
		Non-English language studies
11.	Context	Although not in the guideline Scope document it was agreed to be important because there is currently variation in practice. Also, pharmacological management in asymptomatic HVD can be used with the aim of preventing the development of heart failure as consequence of heart valve disease, so the impact is clinically relevant. Statins and alpha-blockers are relevant to the asymptomatic population but not those with concomitant heart failure and so have been added to this protocol although not listed in the Scope.
12.	Primary outcomes (critical outcomes)	<ul> <li>All-cause mortality at ≥12 months (dichotomous)</li> </ul>
		• Cardiac mortality at ≥12 months (dichotomous)
		<ul> <li>Health-related quality of life at 6 months and ≥12 months (continuous)</li> </ul>
		<ul> <li>Onset of symptoms or progression in NYHA class at ≥12 months</li> </ul>
		<ul> <li>Evidence of HVD progression on imaging (worsening of disease severity) at ≥ 12 months (dichotomous)</li> </ul>
		• Need for heart valve intervention (surgical or transcatheter) at ≥12 months (dichotomous)

		Follow-up:
		<ul> <li>Include only the closest reported time to the 6 month time-point from each study if multiple time points are recorded.</li> </ul>
		<ul> <li>Report the longest follow-up time reported for the ≥12 month time-point if multiple time-points are recorded</li> </ul>
		No minimum time-point for inclusion
13.	Secondary outcomes (important outcomes)	<ul> <li>Exercise tolerance reported as any of the following (in order of relevance) at 12 months:         <ul> <li>Supine bicycle workload (watts or % difference from predicted watts)</li> <li>Treadmill exercise time (duration)</li> <li>Oxygen consumption on exercise testing (VO₂ max)</li> <li>Time to near maximal dyspnoea</li> <li>6-minute walk test</li> <li>Borg dyspnoea index</li> </ul> </li> <li>(Continuous, final values or change scores – choose the type most often reported in other studies if both available in a single study, combine change and final scores in meta-analysis if appropriate)</li> <li>Withdrawal from the trial due to adverse events at 6 and 12 months (dichotomous)</li> <li>Follow-up:         <ul> <li>Include only the closest reported time to the 6 month time-point from each study if multiple time points are recorded.</li> <li>Report the longest follow-up time reported for the ≥12 month time-point if multiple time-points are recorded</li> <li>No minimum time-point for inclusion</li> </ul> </li> </ul>
14.	Data extraction (selection and coding)	EndNote will be used for reference management, sifting, citations and bibliographies. All references identified by the searches and from other sources will be screened for inclusion. 10% of the abstracts will be reviewed by two reviewers, with any disagreements resolved by discussion or, if necessary, a third independent reviewer.
		The full text of potentially eligible studies will be retrieved and will be assessed in line with the criteria outlined above.

		An in-house developed database; EviBase, will be used for data extraction. A standardised form is followed to extract data from studies (see Developing NICE guidelines: the manual section 6.4) and for undertaking assessment of study quality. Summary evidence tables will be produced including information on: study setting; study population and participant demographics and baseline characteristics; details of the intervention and control interventions; study methodology' recruitment and missing data rates; outcomes and times of measurement; critical appraisal ratings.
		MS Excel will be used for data extraction and critical appraisal for health economic studies.
15.	Risk of bias (quality) assessment	Risk of bias will be assessed using the appropriate checklist as described in Developing NICE guidelines: the manual.
		Checklists used in this intervention review are as follows for different types of study design:
		Systematic reviews: Risk of Bias in Systematic Reviews (ROBIS)
		Randomised Controlled Trial: Cochrane RoB (2.0)
		10% of all evidence reviews are quality assured by a senior research fellow. This includes checking:
		papers were included /excluded appropriately
		a sample of the data extractions
		correct methods are used to synthesise data
		a sample of the risk of bias assessments
		Disagreements between the review authors over the risk of bias in particular studies will be resolved by discussion, with involvement of a third party where necessary.
16.	Strategy for data synthesis	Where possible, data will be meta-analysed. Pairwise meta-analyses will be performed using Cochrane Review Manager (RevMan5) to combine the data given in all studies for each of the outcomes stated above. A fixed effect meta-analysis, with weighted mean differences for continuous outcomes and risk ratios for binary outcomes will be used, and 95% confidence intervals will be calculated for each outcome.
		<ul> <li>Heterogeneity between the studies in effect measures will be assessed using the I² statistic and visually inspected. We will consider an I² value greater than 50% indicative of substantial heterogeneity. Sensitivity analyses will be conducted based on pre-specified subgroups using stratified meta-analysis to explore the heterogeneity in effect estimates. If this does not explain the heterogeneity, the results will be presented using random-effects.</li> </ul>
		GRADEpro will be used to assess the quality of evidence for each outcome, taking into account individual study quality and the meta-analysis results. The 4 main quality elements (risk of bias,

		<ul> <li>indirectness, inconsistency and imprecision) will be appraised for each outcome. Publication bias is tested for when there are more than 5 studies for an outcome.</li> <li>WinBUGS will be used for network meta-analysis, if possible given the data identified.</li> <li>Where meta-analysis is not possible, data will be presented and quality assessed individually per outcome.</li> <li>A second reviewer will quality assure 10% of the data analyses. Discrepancies will be identified and</li> </ul>	
		resolved through discussion (with a third party where necessary).	
17.	Analysis of sub-groups	<ul> <li>Groups that will be analysed separately (strata):         <ul> <li>Primary aortic [including bicuspid] stenosis</li> <li>Primary aortic regurgitation</li> <li>Primary mitral stenosis</li> <li>Primary mitral regurgitation</li> <li>Primary tricuspid regurgitation</li> <li>Secondary heart valve disease – mitral regurgitation or tricuspid regurgitation</li> </ul> </li> <li>Subgroups that will be investigated if heterogeneity is present:         <ul> <li>Presence vs. absence of uncontrolled systemic hypertension (140/85 mmHg) at the end of trial intervention despite medication</li> <li>Age (&lt;75 vs. ≥75 years)</li> <li>Disease mechanism:             <ul> <li>Aortic and mitral stenosis: calcific vs non-calcific</li> </ul> </li> </ul></li></ul>	
18.	Type and method of review	☑ Intervention   ☐ Diagnostic   ☐ Prognostic   ☐ Qualitative   ☐ Epidemiologic   ☐ Service Delivery   ☐ Other (please specify)	

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19.	Language	English				
20.	Country	England				
21.	Anticipated or actual start date	09/05/2019				
22.	Anticipated completion date	17/06/2021				
23.	Stage of review at time of this submission	Review stage	Started	Completed		
		Preliminary searches	~			
		Piloting of the study selection process	<b>V</b>			
		Formal screening of search results against eligibility criteria	~			
		Data extraction	~			
		Risk of bias (quality) assessment	V			
		Data analysis	•	<u> </u>		
24.	Named contact	5a. Named contact National Guideline Centre				
		5b Named contact e-mail				
		HVD@nice.org.uk				
		5e Organisational affiliation of the review				
		National Institute for Healt	h and Care Ex	xcellence (NICE) and the National Guideline Centre		
25.	Review team members	From the National Guideline Centre:				
	Sharon Swain [Guideline lead]					

		Eleanor Samarasekera [Senior systematic reviewer]		
		Nicole Downes [Systematic reviewer]		
		George Wood [Systematic reviewer]		
		Robert King [Health economist]		
		Jill Cobb [Information specialist]		
		Claire Townsend [Information specialist]		
		Katie Broomfield [Project manager]		
26.	Funding sources/sponsor	This systematic review is being completed by the National Guideline Centre which receives funding from NICE.		
27.	Conflicts of interest	All guideline committee members and anyone who has direct input into NICE guidelines (including the evidence review team and expert witnesses) must declare any potential conflicts of interest in line with NICE's code of practice for declaring and dealing with conflicts of interest. Any relevant interests, or changes to interests, will also be declared publicly at the start of each guideline committee meeting. Before each meeting, any potential conflicts of interest will be considered by the guideline committee Chair and a senior member of the development team. Any decisions to exclude a person from all or part of a meeting will be documented. Any changes to a member's declaration of interests will be recorded in the minutes of the meeting. Declarations of interests will be published with the final guideline.		
28.	Collaborators	Development of this systematic review will be overseen by an advisory committee who will use the review to inform the development of evidence-based recommendations in line with section 3 of <a href="Developing NICE guidelines: the manual">Developing NICE guidelines: the manual</a> . Members of the guideline committee are available on the NICE website: <a href="https://www.nice.org.uk/guidance/indevelopment/gid-ng10122">https://www.nice.org.uk/guidance/indevelopment/gid-ng10122</a>		
29.	Other registration details	N/A		
30.	Reference/URL for published protocol	_		
31.	Dissemination plans	NICE may use a range of different methods to raise awareness of the guideline. These include standard approaches such as:		
		notifying registered stakeholders of publication		
		publicising the guideline through NICE's newsletter and alerts		
		• issuing a press release or briefing as appropriate, posting news articles on the NICE website, using social media channels, and publicising the guideline within NICE.		

32.	Keywords	regurgitation; inhibitors; an	Heart valve disease; pharmacological treatment; medical treatment; asymptomatic; aortic stenosis; aortic regurgitation; mitral stenosis; mitral regurgitation; tricuspid regurgitation; angiotensin-converting enzyme inhibitors; angiotensin II receptor antagonists; beta blockers; calcium channel blockers; digoxin; diuretics; statins; nitrates.		
33.	Details of existing review of same topic by same authors	N/A	·		
34.	Current review status		Ongoing		
		$\boxtimes$	Completed but not published		
			Completed and published		
			Completed, published and being updated		
			Discontinued		
35.	Additional information	N/A			
36.	Details of final publication	www.nice.org.uk			

# A.2 Valve disease with heart failure

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4 Table 27: Review protocol: pharmacological management with heart failure

ID	Field	Content
0.	PROSPERO registration number	
1.	Review title	In adults with heart failure and heart valve disease, what is the clinical and cost effectiveness of angiotensin-converting enzyme (ACE) inhibitors, angiotensin-II receptor blockers (ARBs), beta blockers, calcium channel blockers, digoxin, diuretics and nitrates to improve clinical outcome?
2.	Review question	In adults with heart failure and concomitant heart valve disease, what is the clinical and cost effectiveness of ACE inhibitors, ARBs, beta blockers, calcium channel blockers, digoxin, diuretics and nitrates to improve clinical outcome?

ID	Field	Content	
3.	Objective	To assess the clinical and cost-effectiveness of pharmacological interventions individually and in combination to manage heart failure specifically in adults with heart valve disease.	
4.	Searches	The following databases will be searched: Cochrane Central Register of Controlled Trials (CENTRAL) Cochrane Database of Systematic Reviews (CDSR) Embase MEDLINE	
		Searches will be restricted by:	
		English language studies	
		Human studies	
		Letters and comments are excluded	
		Validated study filters for systematic reviews and RCTs	
		No date restrictions applied	
		The searches may be re-run 6 weeks before final committee meeting and further studies retrieved for inclusion if relevant.	
		The full search strategies database will be published in the final review.	
5.	Condition or domain being studied	Diagnosed heart valve disease in adults aged 18 years and over: Aortic (including bicuspid) stenosis, aortic regurgitation, mitral stenosis, mitral regurgitation and tricuspid regurgitation.	
6.	Population	Inclusion:  Adults aged 18 years and over with diagnosed heart failure and heart valve disease of at least moderate severity stratified by type:  Primary aortic [including bicuspid] stenosis  Primary aortic regurgitation  Primary mitral stenosis  Primary mitral regurgitation  Primary tricuspid regurgitation  Secondary heart valve disease – mitral regurgitation or tricuspid regurgitation	

ID	Field	Content
		Any heart failure definition will be accepted as reported in the studies, with downgrading for risk of bias/indirectness if not adequately or appropriately defined. As stated in the NICE guideline on chronic heart failure (NG106), it is a complex clinical syndrome of symptoms and signs caused by impairment of the heart's action as a pump supporting the circulation. It is caused by structural or functional abnormalities of the heart. The demonstration of objective evidence of these cardiac abnormalities is necessary for the diagnosis of heart failure to be made. The symptoms most commonly encountered are breathlessness (exertional dyspnoea, orthopnoea and paroxysmal nocturnal dyspnoea), fatigue, and oedema. Signs in heart failure could be due to pulmonary and systemic congestion, or the structural abnormalities either causing or caused by heart failure. Therefore, the ideal definition includes symptoms and/or signs plus associated structural or functional abnormalities that can explain the symptoms and signs.  Heart valve disease severity should be reported according to standard thresholds from echocardiography, as reported by the British Society of Echocardiography. However, study definitions will be accepted and discussed with the committee to determine whether they represent indirect evidence. If severity is not stated, the study will be included but downgraded for indirectness.
		Include both those with no current plan/need for intervention and those who are receiving drugs as bridging therapy while waiting for an intervention. These groups will be pooled initially but considered for subgroup analysis if heterogeneity is found.
		Include only first line use of pharmacological management options.
		In cases of mixed heart valve disease (i.e. the patients each had more than one type of valve disease) the study will be classified according to the predominant valve lesion that drives medical decision making (e.g. in rheumatic heart disease, this may be mitral stenosis).
		Studies including adults with HVD but not all with concomitant heart failure will be included if ≥50% of the included patients had heart failure. Those with <50% having HF will be included in a separate question on HVD without heart failure.
		Groups from the equality impact assessment were considered. It was decided that they did not need to be considered separately for this question,

ID	Field	Content
		Inclusion of indirect evidence from mixed populations, to be considered separately for each strata and intervention:
		If no/insufficient studies are found in the HVD population, studies including adults with heart failure from mixed causes will be included if >75% of the included patients had HVD.
		Exclusions:
		Pharmacological management in children (17 years and under)
		People with congenital heart valve disease, except bicuspid aortic valve disease.
7.	Intervention/Exposure/Test	Angiotensin-converting enzyme (ACE) inhibitors
		Angiotensin-II receptor blockers (ARBs)
		Beta blockers Calcium channel blockers
		Digoxin
		Diuretics
		Nitrates (including nitroprusside)
		Any combination of 2 or more of the above
		Primary studies with a mixed intervention (some in the 'active' arm received the intervention of interest and some a different intervention) will be included if at least 90% received the intervention of interest.
		A class effect will be used for analysis, combining all interventions within each drug class (regardless of mode of delivery, and dose – as long as within the licensed range). Based on this class effect assumption, drugs not licenced for HF within a class that has some licenced agents could be considered as long as not contraindicated.
		For crossover studies, there is no lower limit for the washout period because drug effects do not persist once a person is no longer taking the drug.
8.	Comparator/Reference	Placebo or no treatment
	standard/Confounding factors	Usual care (e.g. following standard heart failure guidelines: ACE + beta-blocker + diuretic) Other active comparator listed above, including combinations
9.	Types of study to be included	Randomised control trials (RCTs) or systematic reviews of RCTs, including crossover trials

ID	Field	Content		
		If no RCT data is available, observational data will not be considered for pharmacological interventions. This is due to the risk of confounding variables influencing the study results, reducing our confidence in the review results.		
10.	Other exclusion criteria	Exclusion criteria:  Conference abstracts will be excluded because they are unlikely to contain enough information to assess whether the population matches the review question in terms of previous medication use, or enough detail on outcome definitions, or on the methodology to assess the risk of bias of the study.  Non-randomised studies / observational studies  Pharmacological management of heart failure in adults without HVD  Pharmacological management of mild HVD with associated heart failure  Pharmacological management in the intra- or post-operative period  Studies combining people with different types of valve disease (e.g. some with mitral regurgitation and some with aortic regurgitation).  Non-English language studies		
11.	Context	This is important because there is uncertainty about the most appropriate medicines for pharmacological management, and whether this differs from the guidelines for heart failure not associated with HVD. There is also variation in practice.		
12.	Primary outcomes (critical outcomes)	All-cause mortality at 12 months (dichotomous) Cardiac mortality at 12 months (dichotomous) Hospital admission due to heart failure at 12 months (dichotomous) Health-related quality of life at 6 months and 12 months (continuous)  Follow-up: include only the closest reported time to the 12- or 6-month time-points from each study if multiple time points are recorded no minimum time point for inclusion		
13.	Secondary outcomes (important outcomes)	Exercise tolerance reported as any of the following (in order of relevance):  Treadmill exercise time (duration)  Time to near maximal dyspnoea  6-minute walk test  Borg dyspnoea index  (continuous, final values or change scores – choose the type most often reported in other studies if both available in a single study, combine change and final scores in meta-analysis if appropriate)  Need for heart valve intervention (surgical or transcatheter) within 12 months (dichotomous)		

ID	Field	Content				
		Withdrawal from the study due to adverse events at 6 months and 12 months (dichotomous)				
		Follow-up:				
		include only the closest reported time to the 12- or 6-month time-points from each study if multiple time points are recorded				
		no minimum time point for inclusion				
14.	Data extraction (selection and coding)	EndNote will be used for reference management, sifting, citations and bibliographies. All references identified by the searches and from other sources will be screened for inclusion.				
		The full text of potentially eligible studies will be retrieved and will be assessed in line with the criteria outlined above.				
		An in-house developed database; EviBase, will be used for data extraction. A standardised form is followed to extract data from studies (see Developing NICE guidelines: the manual section 6.4) and for undertaking assessment of study quality. Summary evidence tables will be produced including information on: study setting; study population and participant demographics and baseline characteristics; details of the intervention and control interventions; study methodology' recruitment and missing data rates; outcomes and times of measurement; critical appraisal ratings.				
		10% of the sifting and extractions will be reviewed by two reviewers, with any disagreements resolved by discussion or, if necessary, a third party.				
		MS Excel will be used for data extraction and critical appraisal for health economic studies.				
15.	Risk of bias (quality) assessment	Risk of bias will be assessed using the appropriate checklist as described in Developing NICE guidelines: the manual.				
		Checklists used in this intervention review are as follows for different types of study design: Systematic reviews: Risk of Bias in Systematic Reviews (ROBIS)				
		Randomised Controlled Trial: Cochrane RoB (2.0)				
		A 10% sample of the risk of bias assessments will be independently quality assured by a second reviewer. Disagreements between the review authors over the risk of bias in particular studies will be resolved by discussion, with involvement of a third party where necessary.				
16.	Strategy for data synthesis	Where possible, data will be meta-analysed. Pairwise meta-analyses will be performed using Cochrane Review Manager (RevMan5) to combine the data given in all studies for each of the outcomes stated above. A fixed effect meta-analysis, with weighted mean differences for continuous outcomes and risk ratios for binary outcomes will be used, and 95% confidence intervals will be calculated for each outcome. Heterogeneity between the studies in effect measures will be assessed using the I² statistic and visually inspected. We will consider an I² value greater than 50% indicative of substantial heterogeneity. Sensitivity				
		analyses will be conducted based on pre-specified subgroups using stratified meta-analysis to explore the				

ID	Field	Content					
		heterogeneity in effect estimates. If this does not explain the heterogeneity, the results will be presented using random-effects.					
		GRADEpro will be used to assess the quality of evidence for each outcome, taking into account individual study quality and the meta-analysis results. The 4 main quality elements (risk of bias, indirectness, inconsistency and imprecision) will be appraised for each outcome. Publication bias is tested for when there are more than 5 studies for an outcome.					
		WinBUGS will be used for network meta-analysis, if possible given the data identified.					
		Where meta-analysis is not possible, data will be presented and quality assessed individually per outcome.					
		A second reviewer will quality assure 10% of the data analyses. Discrepancies will be identified and resolved through discussion (with a third party where necessary).					
17.	Analysis of sub-groups	Groups that will be analysed separately (strata)					
		Type of HVD:					
		aortic [including bicuspid] stenosis					
		aortic regurgitation mitral stenosis					
		mitral regurgitation					
		tricuspid regurgitation					
		Cub argume that will be investigated if between partity is present.					
		Subgroups that will be investigated if heterogeneity is present:  Severe vs moderate HVD (as defined by the British Society of Echocardiography)					
		Symptomatic vs asymptomatic					
		Age (<75 versus ≥75)					
		Disease mechanism:					
		Aortic and mitral stenosis: calcific vs non-calcific					
		Presence vs. absence of uncontrolled systemic hypertension (140/85 mmHg) at the end of trial intervention despite medication					
18.	Type and method of review						
		□ Diagnostic					
		□ Prognostic					
		□ Qualitative					
		□ Epidemiologic					

ID	Field	Content					
		□ Service Delivery					
			lease specify)				
19.	Language	English					
20.	Country	England					
21.	Anticipated or actual start date	09/05/2019					
22.	Anticipated completion date	17/06/2021					
23.	Stage of review at time of this	Review stag	ge	Started	Cor	mpleted	
	submission	Preliminary searches		•	~		
		Piloting of the study selection process		•	~		
		Formal screening of search results against eligibility criteria		•	~		
		Data extraction		<b>✓</b>	~		
		Risk of bias (quality) assessment		•	~		
		Data analysis		<b>✓</b>	~		
24.	Named contact	<ul> <li>5a. Named contact</li> <li>National Guideline Centre</li> <li>5b Named contact e-mail</li> <li>HVD@nice.org.uk</li> <li>5e Organisational affiliation of the review</li> </ul>					
		National Institute for Health and Care Excellence (NICE) and the National Guideline Centre					
25.	Review team members	From the National Guideline Centre: Sharon Swain [Guideline lead]					

ID	Field	Content	
		Eleanor Samarasekera [Senior systematic reviewer] Nicole Downes [Systematic reviewer] George Wood [Systematic reviewer] Robert King [Health economist] Jill Cobb [Information specialist] Katie Broomfield [Project manager]	
26.	Funding sources/sponsor	This systematic review is being completed by the National Guideline Centre which receives funding from NICE.	
27.	Conflicts of interest	All guideline committee members and anyone who has direct input into NICE guidelines (including the evidence review team and expert witnesses) must declare any potential conflicts of interest in line with NICE's code of practice for declaring and dealing with conflicts of interest. Any relevant interests, or changes to interests, will also be declared publicly at the start of each guideline committee meeting. Before each meeting, any potential conflicts of interest will be considered by the guideline committee Chair and a senior member of the development team. Any decisions to exclude a person from all or part of a meeting will be documented. Any changes to a member's declaration of interests will be recorded in the minutes of the meeting. Declarations of interests will be published with the final guideline.	
28.	Collaborators	Development of this systematic review will be overseen by an advisory committee who will use the review to inform the development of evidence-based recommendations in line with section 3 of Developing NICE guidelines: the manual. Members of the guideline committee are available on the NICE website: https://www.nice.org.uk/guidance/indevelopment/gid-ng10122	
29.	Other registration details	None	
30.	Reference/URL for published protocol		
31.	Dissemination plans	NICE may use a range of different methods to raise awareness of the guideline. These include standard approaches such as: notifying registered stakeholders of publication publicising the guideline through NICE's newsletter and alerts issuing a press release or briefing as appropriate, posting news articles on the NICE website, using social media channels, and publicising the guideline within NICE.	
32.	Keywords	Heart valve disease; Anticoagulation; Antiplatelet; Biological heart valve; Intervention; Surgical valve replacement; Transcatheter valve replacement	
33.	Details of existing review of same topic by same authors	N/A	

ID	Field	Content		
34.	34. Current review status		Ongoing	
			Completed but not published	
			Completed and published	
			Completed, published and being updated	
			Discontinued	
35.	Additional information	N/A		
36.	Details of final publication	www.nice.org.uk		

# Table 28: Health economic review protocol

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Review question	All questions – health economic evidence					
Objectives	To identify health economic studies relevant to any of the review questions.					
Search criteria	• Populations, interventions and comparators must be as specified in the clinical review protocol above.					
	• Studies must be of a relevant health economic study design (cost-utility analysis, cost-effectiveness analysis, cost-benefit analysis, cost-consequences analysis, comparative cost analysis).					
	• Studies must not be a letter, editorial or commentary, or a review of health economic evaluations. (Recent reviews will be ordered although not reviewed. The bibliographies will be checked for relevant studies, which will then be ordered.)					
	• Unpublished reports will not be considered unless submitted as part of a call for evidence.					
	Studies must be in English.					
Search strategy	A health economic study search will be undertaken using population-specific terms and a health economic study filter – see appendix B below.					
Review strategy	Studies not meeting any of the search criteria above will be excluded. Studies published before 2004, abstract-only studies and studies from non-OECD countries or the USA will also be excluded.					
	Each remaining study will be assessed for applicability and methodological limitations using the NICE economic evaluation checklist which can be found in appendix H of Developing NICE guidelines: the manual (2014). <sup>112</sup>					
	Inclusion and exclusion criteria					
	• If a study is rated as both 'Directly applicable' and with 'Minor limitations' then it will be included in the guideline. A health economic evidence table will be completed and it will be included in the health economic evidence profile.					

- If a study is rated as either 'Not applicable' or with 'Very serious limitations' then it will usually be excluded from the guideline. If it is excluded then a health economic evidence table will not be completed and it will not be included in the health economic evidence profile.
- If a study is rated as 'Partially applicable', with 'Potentially serious limitations' or both then there is discretion over whether it should be included.

#### Where there is discretion

The health economist will make a decision based on the relative applicability and quality of the available evidence for that question, in discussion with the guideline committee if required. The ultimate aim is to include health economic studies that are helpful for decision-making in the context of the guideline and the current NHS setting. If several studies are considered of sufficiently high applicability and methodological quality that they could all be included, then the health economist, in discussion with the committee if required, may decide to include only the most applicable studies and to selectively exclude the remaining studies. All studies excluded on the basis of applicability or methodological limitations will be listed with explanation in the excluded health economic studies appendix below.

The health economist will be guided by the following hierarchies.

#### Setting:

- UK NHS (most applicable).
- OECD countries with predominantly public health insurance systems (for example, France, Germany, Sweden).
- OECD countries with predominantly private health insurance systems (for example, Switzerland).
- Studies set in non-OECD countries or in the USA will be excluded before being assessed for applicability and methodological limitations.

### Health economic study type:

- Cost-utility analysis (most applicable).
- Other type of full economic evaluation (cost-benefit analysis, cost-effectiveness analysis, cost-consequences analysis).
- Comparative cost analysis.
- Non-comparative cost analyses including cost-of-illness studies will be excluded before being assessed for applicability and methodological limitations.

### Year of analysis:

- The more recent the study, the more applicable it will be.
- Studies published in 2004 or later that depend on unit costs and resource data entirely or predominantly from before 2004 will be rated as 'Not applicable'.
- Studies published before 2004 will be excluded before being assessed for applicability and methodological limitations.

Quality and relevance of effectiveness data used in the health economic analysis:

• The more closely the clinical effectiveness data used in the health economic analysis match with the outcomes of the studies included in the clinical review the more useful the analysis will be for decision-making in the guideline.

Appendix B: Literature search strategies

## B.4 Valve disease without heart failure

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- 5 <u>Heart valve disease search strategy 6 pharmacological management without heart failure</u>
- 6 This literature search strategy was used for the following reviews:
  - In adults with heart valve disease without concomitant heart failure, what is the clinical and cost effectiveness of alpha-blockers, angiotensin-converting enzyme (ACE) inhibitors, angiotensin-II receptor blockers (ARBs), beta blockers, calcium channel blockers, digoxin, diuretics, statins and nitrates to improve clinical outcome?
- 11 The literature searches for this review are detailed below and complied with the methodology outlined in Developing NICE guidelines: the manual. 112
- For more information, please see the Methodology review published as part of the accompanying documents for this guideline.

# B.1.11 Clinical search literature search strategy

- 2 Searches were constructed using a PICO framework where population (P) terms were
- 3 combined with Intervention (I) and in some cases Comparison (C) terms. Outcomes (O) are
- 4 rarely used in search strategies for interventions as these concepts may not be well
- 5 described in title, abstract or indexes and therefore difficult to retrieve. Search filters were
- 6 applied to the search where appropriate.

7

# 8 Table 29: Database date parameters and filters used

Database	Dates searched	Search filter used
Medline (OVID)	1946 - 14 October 2020	Exclusions Randomised controlled trials Systematic review studies
Embase (OVID)	1974 - 14 October 2020	Exclusions Randomised controlled trials Systematic review studies
The Cochrane Library (Wiley)	Cochrane Reviews to 2020 Issue 10 of 12 CENTRAL to 2020 Issue 10 of 12	None

#### 9 Medline (Ovid) search terms

<ol> <li>exp heart valves/</li> <li>((primary or secondary) adj valv* disease*).ti,ab.</li> <li>((valv* or flap* or leaflet*) adj1 (heart or cardiac) adj (disease* or disorder* of failed or dysfunction* or insufficien* or repair* or replace* or damage* or leaflet or disorder* or failure or failed or pulmon*) adj (valv* or flap* or leaflet*) adj (disorder* or failure or failed or dysfunction* or insufficien* or repair* or replated amage* or leak*)).ti,ab.</li> <li>((mitral or aortic or tricuspid or pulmon*) adj3 (prolapse or regurgitation or satresia or insufficienc*)).ti,ab.</li> <li>Heart Valve Prosthesis/</li> <li>((mechanical or artificial or prosthe* or bioprosthe* or biological or tissue) a flap* or leaflet*)).ti,ab.</li> <li>valve-in-valve.ti,ab.</li> </ol>	
<ol> <li>((valv* or flap* or leaflet*) adj1 (heart or cardiac) adj (disease* or disorder* of failed or dysfunction* or insufficien* or repair* or replace* or damage* or leaflet</li> <li>((mitral or aortic or tricuspid or pulmon*) adj (valv* or flap* or leaflet*) adj (disorder* or failure or failed or dysfunction* or insufficien* or repair* or replated amage* or leak*)).ti,ab.</li> <li>((mitral or aortic or tricuspid or pulmon*) adj3 (prolapse or regurgitation or statresia or insufficienc*)).ti,ab.</li> <li>Heart Valve Prosthesis/</li> <li>((mechanical or artificial or prosthe* or bioprosthe* or biological or tissue) at flap* or leaflet*)).ti,ab.</li> <li>valve-in-valve.ti,ab.</li> </ol>	
failed or dysfunction* or insufficien* or repair* or replace* or damage* or leaflet  ((mitral or aortic or tricuspid or pulmon*) adj (valv* or flap* or leaflet*) adj (d disorder* or failure or failed or dysfunction* or insufficien* or repair* or replated damage* or leak*)).ti,ab.  ((mitral or aortic or tricuspid or pulmon*) adj3 (prolapse or regurgitation or satresia or insufficienc*)).ti,ab.  Heart Valve Prosthesis/  ((mechanical or artificial or prosthe* or bioprosthe* or biological or tissue) a flap* or leaflet*)).ti,ab.	
disorder* or failure or failed or dysfunction* or insufficien* or repair* or replated amage* or leak*)).ti,ab.  ((mitral or aortic or tricuspid or pulmon*) adj3 (prolapse or regurgitation or satresia or insufficienc*)).ti,ab.  Heart Valve Prosthesis/  ((mechanical or artificial or prosthe* or bioprosthe* or biological or tissue) a flap* or leaflet*)).ti,ab.  valve-in-valve.ti,ab.	
atresia or insufficienc*)).ti,ab.  7. Heart Valve Prosthesis/  8. ((mechanical or artificial or prosthe* or bioprosthe* or biological or tissue) a flap* or leaflet*)).ti,ab.  9. valve-in-valve.ti,ab.	
<ul> <li>8. ((mechanical or artificial or prosthe* or bioprosthe* or biological or tissue) a flap* or leaflet*)).ti,ab.</li> <li>9. valve-in-valve.ti,ab.</li> </ul>	stenos?s or
flap* or leaflet*)).ti,ab.  9. valve-in-valve.ti,ab.	
	dj (valv* or
10. (transcatheter adj2 (valve or valves)).ti,ab.	
11. exp Heart Murmurs/	
12. ((heart or cardiac) adj murmur*).ti,ab.	
13. or/1-12	
14. letter/	
15. editorial/	
16. news/	
17. exp historical article/	
18. Anecdotes as Topic/	
19. comment/	
20. case report/	
21. (letter or comment*).ti.	
22. or/14-21	

<sup>&</sup>lt;Click this field on the first page and insert footer text if required>
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23.	randomized controlled trial/ or random*.ti,ab.	
24.	22 not 23	
25.	animals/ not humans/	
26.	exp Animals, Laboratory/	
27.	exp Animals, East-activities exp Animal Experimentation/	
28.	exp Models, Animal/	
29.	exp Rodentia/	
30.	(rat or rats or mouse or mice).ti.	
31.	or/24-30	
32.	13 not 31	
33.	limit 32 to English language	
34.	(exp child/ or exp pediatrics/ or exp infant/) not (exp adolescent/ or exp adult/ or exp middle age/ or exp aged/)	
35.	33 not 34	
36.	randomized controlled trial.pt.	
37.	controlled clinical trial.pt.	
38.	randomi#ed.ti,ab.	
39.	placebo.ab.	
40.	randomly.ti,ab.	
41.	Clinical Trials as topic.sh.	
42.	trial.ti.	
43.	or/36-42	
44.	Meta-Analysis/	
45.	exp Meta-Analysis as Topic/	
46.	(meta analy* or metanaly* or meta regression).ti,ab.	
47.	((systematic* or evidence*) adj3 (review* or overview*)).ti,ab.	
48.	(reference list* or bibliograph* or hand search* or manual search* or relevant journals).ab.	
49.	(search strategy or search criteria or systematic search or study selection or data extraction).ab.	
50.	(search* adj4 literature).ab.	
51.	(medline or pubmed or cochrane or embase or psychlit or psyclit or psychinfo or psycinfo or cinahl or science citation index or bids or cancerlit).ab.	
52.	cochrane.jw.	
53.	((multiple treatment* or indirect or mixed) adj2 comparison*).ti,ab.	
54.	or/44-52	
55.	35 and (43 or 54)	
56.	exp Angiotensin-Converting Enzyme Inhibitors/	
57.	(("angiotensin-converting enzyme" or ace) adj2 (inhibitor* or antagonist*)).ti,ab.	
58.	(Captopril or cilazapril or enalapril or enalaprilat or fosinopril or lisinopril or perindopril or quinapril or ramipril or teprotide or imidapril or trandolopril).ti,ab.	
59.	exp Angiotensin Receptor Antagonists/	
60.	(angiotensin adj3 receptor adj3 (antagonist* or blocker*)).ti,ab.	
61.	(1-Sarcosine-8-Isoleucine or amlodipine or irbesartan or olmesartan or saralasin or telmisartan or valsartan or azilsartan or candesartan or eprosartan or losartan).ti,ab.	
62.	exp Adrenergic beta-Antagonists/	

63.	(Acebutolol or Atenolol or Bisoprolol or Carvedilol or Celiprolol or Esmolol or Labetalol or Metoprolol or Nadolol or Nebivolol or Oxprenolol or Propranolol or Pindolol or Sotalol or Timolol).ti,ab.
64.	(beta adj3 block*).ti,ab.
65.	((beta-adrenoceptor or b-adrenoceptor or beta-adrenergic) adj (block* or antagonist*)).ti,ab.
66.	(b adj3 block*).ti,ab.
67.	(beta adj2 antagonist*).ti,ab.
68.	exp Calcium Channel Blockers/
69.	(calcium adj3 (block* or antagonis* or inhibit*)).ti,ab.
70.	calcium channel receptor block*.ti,ab.
71.	(amlodipine or amrinone or bencyclane or bepridil or carvedilol or cinnarizine or conotoxin* or diltiazem or felodipine or fendiline or flunarizine or gallopamil or isradipine or lamotrigine or mibefradil or nicardipine or nifedipine or nimodipine or nisoldipine or nitrendipine or perhexiline or pregabalin or prenylamine or risedronic acid or tiapamil or verapamil or zonisamide or atenolol or clevidipine or lacidipine or lercanidipine or dihyropyridine*).ti,ab.
72.	exp Digoxin/
73.	digoxin.ti,ab.
74.	exp Diuretics/
75.	((oral* or subcut* or IV or intravenous* or iv or infusion* or drip or drips or augment* or sequential* or loop or "high ceiling") adj6 diuretic*).ti,ab.
76.	(augment* adj diuresis).ti,ab.
77.	(acetazolamide or amiloride or bendroflumenethiazide or bumentanide or chlorothiazide or chlorothalidone or clopamide or cyclopenhiazide or ethacrynic acid or ethoxzolamide or furosemide or hydrochlorothiazide or hydroflumethiazide or indapamide or mefruside or methazolamide or methyclothiazide or metolazone or muzolimine or polythiazide or potassium citrate or spironolactone or ticrynafen or torsemide or triamterene or trichlormethiazide or xipamide or isosorbide or mannitol or co-amilozide or co-triamterzide or co-flumactone or eplerenone or co-amilofruse).ti,ab.
78.	exp Nitrates/
79.	Nitroglycerin/
80.	Nitroprusside/
81.	Isosorbide Dinitrate/
82.	(nitroglycerin or gtn or nitrate* or nitroprusside*).ti,ab.
83.	(glyceryl trinitrate or isosorbide dinitrate or isosorbide mononitrate).ti,ab.
84.	*hydroxymethylglutaryl-coa reductase inhibitors/ or atorvastatin calcium/ or pravastatin/ or rosuvastatin calcium/ or exp simvastatin/
85.	(atorvastatin* or pravastatin* or rosuvastatin* or simvastatin* or fluvastatin*).ti,ab.
86.	((Hydroxymethylglutaryl-CoA or HMG-CoA or Hydroxymethylglutaryl-Coenzyme A) adj3 (reductase* or inhibitor*)).ti,ab.
87.	statin*.ti,ab.
88.	exp Adrenergic alpha-Antagonists/
89.	(alfuzosin or bunazosin or doxazosin or indoramin or metazosin or neldazosin or prazosin or silodosin or tamsulosin or terazosin or tiodazosin or trimazosin).ti,ab.
90.	(alpha adrenergic antagonist* or alpha adrenergic receptor antagonist* or adrenergic alpha antagonist*).ti,ab.
91.	((alpha or alpha-adrenergic) adj2 block*).ti,ab.
92.	or/56-91
93.	55 and 92

1 Embase (Ovid) search terms

1.	exp valvular heart disease/
2.	exp heart valve/
3.	((primary or secondary) adj valv* disease*).ti,ab.
	((valv* or flap* or leaflet*) adj1 (heart or cardiac) adj (disease* or disorder* or failure or
4.	failed or dysfunction* or insufficien* or repair* or replace* or damage* or leak*)).ti,ab.
5.	((mitral or aortic or tricuspid or pulmon*) adj (valv* or flap* or leaflet*) adj (disease* or disorder* or failure or failed or dysfunction* or insufficien* or repair* or replace* or damage* or leak*)).ti,ab.
6.	((mitral or aortic or tricuspid or pulmon*) adj3 (prolapse or regurgitation or stenos?s or atresia or insufficienc*)).ti,ab.
7.	exp heart valve prosthesis/
8.	((mechanical or artificial or prosthe* or bioprosthe* or biological or tissue) adj (valv* or flap* or leaflet*)).ti,ab.
9.	valve-in-valve.ti,ab.
10.	(transcatheter adj2 (valve or valves)).ti,ab.
11.	exp heart murmur/
12.	((heart or cardiac) adj murmur*).ti,ab.
13.	or/1-12
14.	letter.pt. or letter/
15.	note.pt.
16.	editorial.pt.
17.	Case report/ or Case study/
18.	(letter or comment*).ti.
19.	or/14-18
20.	randomized controlled trial/ or random*.ti,ab.
21.	19 not 20
22.	animal/ not human/
23.	Nonhuman/
24.	exp Animal Experiment/
25.	exp Experimental animal/
26.	Animal model/
27.	exp Rodent/
28.	(rat or rats or mouse or mice).ti.
29.	or/21-28
30.	13 not 29
31.	limit 30 to English language
32.	(exp child/ or exp pediatrics/) not (exp adult/ or exp adolescent/)
33.	31 not 32
34.	random*.ti,ab.
35.	factorial*.ti,ab.
36.	(crossover* or cross over*).ti,ab.
37.	((doubl* or singl*) adj blind*).ti,ab.
38.	(assign* or allocat* or volunteer* or placebo*).ti,ab.
39.	crossover procedure/
40.	single blind procedure/
41.	randomized controlled trial/

42.	double blind procedure/
43.	or/34-42
44.	systematic review/
45.	meta-analysis/
46.	(meta analy* or metanaly* or metaanaly* or meta regression).ti,ab.
47.	((systematic or evidence) adj3 (review* or overview*)).ti,ab.
48.	(reference list* or bibliograph* or hand search* or manual search* or relevant journals).ab.
49.	(search strategy or search criteria or systematic search or study selection or data extraction).ab.
50.	(search* adj4 literature).ab.
51.	(medline or pubmed or cochrane or embase or psychlit or psyclit or psychinfo or psycinfo or cinahl or science citation index or bids or cancerlit).ab.
52.	((pool* or combined) adj2 (data or trials or studies or results)).ab.
53.	cochrane.jw.
54.	((multiple treatment* or indirect or mixed) adj2 comparison*).ti,ab.
55.	or/44-53
56.	33 and (43 or 55)
57.	exp dipeptidyl carboxypeptidase inhibitor/
58.	(("angiotensin-converting enzyme" or ace) adj2 (inhibitor* or antagonist*)).ti,ab.
59.	(Captopril or cilazapril or enalapril or enalaprilat or fosinopril or lisinopril or perindopril or quinapril or ramipril or teprotide or imidapril or trandolopril).ti,ab.
60.	exp angiotensin receptor antagonist/
61.	(angiotensin adj3 receptor adj3 (antagonist* or blocker*)).ti,ab.
62.	(1-Sarcosine-8-Isoleucine or amlodipine or irbesartan or olmesartan or saralasin or telmisartan or valsartan or azilsartan or candesartan or eprosartan or losartan).ti,ab.
63.	exp beta adrenergic receptor blocking agent/
64.	(Acebutolol or Atenolol or Bisoprolol or Carvedilol or Celiprolol or Esmolol or Labetalol or Metoprolol or Nadolol or Nebivolol or Oxprenolol or Propranolol or Pindolol or Sotalol or Timolol).ti,ab.
65.	(beta adj3 block*).ti,ab.
66.	((beta-adrenoceptor or b-adrenoceptor or beta-adrenergic) adj (block* or antagonist*)).ti,ab.
67.	(b adj3 block*).ti,ab.
68.	(beta adj2 antagonist*).ti,ab.
69.	exp Calcium Channel Blockers/
70.	(calcium adj3 (block* or antagonis* or inhibit*)).ti,ab.
71.	calcium channel receptor block*.ti,ab.
72.	(amlodipine or amrinone or bencyclane or bepridil or carvedilol or cinnarizine or conotoxin* or diltiazem or felodipine or fendiline or flunarizine or gallopamil or isradipine or lamotrigine or mibefradil or nicardipine or nifedipine or nimodipine or nisoldipine or nitrendipine or perhexiline or pregabalin or prenylamine or risedronic acid or tiapamil or verapamil or zonisamide or atenolol or clevidipine or lacidipine or lercanidipine or dihyropyridine*).ti,ab.
73.	digoxin/
74.	digoxin.ti,ab.
75.	exp diuretic agent/
76.	((oral* or subcut* or IV or intravenous* or iv or infusion* or drip or drips or augment* or sequential* or loop or "high ceiling") adj6 diuretic*).ti,ab.

77.	(augment* adj diuresis).ti,ab.
78.	(acetazolamide or amiloride or bendroflumenethiazide or bumentanide or chlorothiazide or chlorthalidone or clopamide or cyclopenhiazide or ethacrynic acid or ethoxzolamide or furosemide or hydrochlorothiazide or hydroflumethiazide or indapamide or mefruside or methazolamide or methyclothiazide or metolazone or muzolimine or polythiazide or potassium citrate or spironolactone or ticrynafen or torsemide or triamterene or trichlormethiazide or xipamide or isosorbide or mannitol or co-amilozide or co-triamterzide or co-flumactone or eplerenone or co-amilofruse).ti,ab.
79.	nitrate/
80.	glyceryl trinitrate/
81.	nitroprusside sodium/
82.	isosorbide mononitrate/
83.	isosorbide dinitrate/
84.	(nitroglycerin or gtn or nitrate* or nitroprusside*).ti,ab.
85.	(glyceryl trinitrate or isosorbide dinitrate or isosorbide mononitrate).ti,ab.
86.	*hydroxymethylglutaryl coenzyme a reductase inhibitor/ or exp atorvastatin/ or exp pravastatin/ or exp.mp. [mp=title, abstract, heading word, drug trade name, original title, device manufacturer, drug manufacturer, device trade name, keyword, floating subheading word, candidate term word]
87.	(atorvastatin* or pravastatin* or rosuvastatin* or simvastatin* or fluvastatin*).ti,ab.
88.	((Hydroxymethylglutaryl-CoA or HMG-CoA or Hydroxymethylglutaryl-Coenzyme A) adj3 (reductase* or inhibitor*)).ti,ab.
89.	statin*.ti,ab.
90.	exp alpha adrenergic receptor blocking agent/
91.	(alfuzosin or bunazosin or doxazosin or indoramin or metazosin or neldazosin or prazosin or silodosin or tamsulosin or terazosin or tiodazosin or trimazosin).ti,ab.
92.	(alpha adrenergic antagonist* or alpha adrenergic receptor antagonist* or adrenergic alpha antagonist*).ti,ab.
93.	((alpha or alpha-adrenergic) adj2 block*).ti,ab.
94.	or/57-93
95.	56 and 94

#### 1 Cochrane Library (Wiley) search terms

#1.	MeSH descriptor: [Heart Valve Diseases] explode all trees
#2.	MeSH descriptor: [Heart Valves] explode all trees
#3.	((primary or secondary) NEXT valv* disease*):ti,ab
#4.	((valv* or flap* or leaflet*) near/2 (heart or cardiac) NEXT (disease* or disorder* or failure or failed or dysfunction* or insufficien* or repair* or replace* or damage* or leak*)):ti,ab
#5.	((mitral or aortic or tricuspid or pulmon*) NEXT (valv* or flap* or leaflet*) NEXT (disease* or disorder* or failure or failed or dysfunction* or insufficien* or repair* or replace* or damage* or leak*)):ti,ab
#6.	((mitral or aortic or tricuspid or pulmon*) NEAR/3 (prolapse or regurgitation or stenos?s or atresia or insufficienc*)):ti,ab
#7.	MeSH descriptor: [Heart Valve Prosthesis] explode all trees
#8.	((mechanical or artificial or prosthe* or bioprosthe* or biological or tissue) NEXT (valv* or flap* or leaflet*)):ti,ab
#9.	valve-in-valve:ti,ab
#10.	(transcatheter NEAR/2 (valve or valves)):ti,ab
#11.	MeSH descriptor: [Heart Murmurs] explode all trees
#12.	((heart or cardiac) NEXT murmur*):ti,ab

#13.	(or #1-#12)	
#14.	MeSH descriptor: [Angiotensin-Converting Enzyme Inhibitors] explode all trees	
#15.	(("angiotensin-converting enzyme" or ace) near/2 (inhibitor* or antagonist*)):ti,ab	
#16.	(Captopril or cilazapril or enalapril or enalaprilat or fosinopril or lisinopril or perindopril or quinapril or ramipril or teprotide or imidapril or trandolopril):ti,ab	
#17.	MeSH descriptor: [Angiotensin Receptor Antagonists] explode all trees	
#18.	(angiotensin near/3 receptor near/3 (antagonist* or blocker*)):ti,ab	
#19.	(amlodipine or irbesartan or olmesartan or saralasin or telmisartan or valsartan or azilsartan or candesartan or eprosartan or losartan):ti,ab	
#20.	1-Sarcosine-8-Isoleucine:ti,ab	
#21.	MeSH descriptor: [Adrenergic beta-Antagonists] explode all trees	
#22.	(Acebutolol or Atenolol or Bisoprolol or Carvedilol or Celiprolol or Esmolol or Labetalol or Metoprolol or Nadolol or Nebivolol or Oxprenolol or Propranolol or Pindolol or Sotalo or Timolol):ti,ab	
#23.	(beta near/3 block*):ti,ab	
#24.	((beta-adrenoceptor or b-adrenoceptor or beta-adrenergic) NEXT (block* or antagonist*)):ti,ab	
#25.	(b near/3 block*):ti,ab	
#26.	(beta near/2 antagonist*):ti,ab	
#27.	MeSH descriptor: [Calcium Channel Blockers] explode all trees	
#28.	(calcium near/3 (block* or antagonis* or inhibit*)):ti,ab	
#29.	'calcium channel receptor':ti,ab	
#30.	(amlodipine or amrinone or bencyclane or bepridil or carvedilol or cinnarizine or conotoxin* or diltiazem or felodipine or fendiline or flunarizine or gallopamil or isradipine or lamotrigine or mibefradil or nicardipine or nifedipine or nimodipine or nisoldipine or nitrendipine or perhexiline or pregabalin or prenylamine or risedronic acid or tiapamil or verapamil or zonisamide or atenolol or clevidipine or lacidipine or lercanidipine or dihyropyridine*):ti,ab	
#31.	MeSH descriptor: [Digoxin] explode all trees	
#32.	digoxin:ti,ab	
#33.	MeSH descriptor: [Diuretics] explode all trees	
#34.	((oral* or subcut* or IV or intravenous* or iv or infusion* or drip or drips or augment* or sequential* or loop or "high ceiling") near/6 diuretic*):ti,ab	
#35.	(acetazolamide or amiloride or bendroflumenethiazide or bumentanide or chlorothiazide or chlorothalidone or clopamide or cyclopenhiazide or ethacrynic acid or ethoxzolamide or furosemide or hydrochlorothiazide or hydroflumethiazide or indapamide or mefruside or methazolamide or methyclothiazide or metolazone or muzolimine or polythiazide or potassium citrate or spironolactone or ticrynafen or torsemide or triamterene or trichlormethiazide or xipamide or isosorbide or mannitol or co-amilozide or co-triamterzide or co-flumactone or eplerenone or co-amilofruse):ti,ab	
#36.	MeSH descriptor: [Nitrates] explode all trees	
#37.	MeSH descriptor: [Nitroglycerin] explode all trees	
#38.	MeSH descriptor: [Nitroprusside] explode all trees	
#39.	MeSH descriptor: [Isosorbide Dinitrate] explode all trees	
#40.	(nitroglycerin or gtn or nitrate* or nitroprusside*):ti,ab	
#41.	('glyceryl trinitrate' or 'isosorbide dinitrate' or 'isosorbide mononitrate'):ti,ab	
#42.	MeSH descriptor: [Hydroxymethylglutaryl-CoA Reductase Inhibitors] explode all trees	
	MeSH descriptor: [Atorvastatin] explode all trees	
#43.		
#43. #44.	MeSH descriptor: [Pravastatin] explode all trees	

#46.	MeSH descriptor: [Simvastatin] explode all trees
#47.	(atorvastatin* or pravastatin* or rosuvastatin* or simvastatin* or fluvastatin*):ti,ab
#48.	((Hydroxymethylglutaryl-CoA or HMG-CoA or Hydroxymethylglutaryl-Coenzyme A) near/3 (reductase* or inhibitor*)):ti,ab
#49.	statin*:ti,ab
#50.	MeSH descriptor: [Adrenergic alpha-Antagonists] explode all trees
#51.	(alfuzosin or bunazosin or doxazosin or indoramin or metazosin or neldazosin or prazosin or silodosin or tamsulosin or terazosin or tiodazosin or trimazosin):ti,ab
#52.	(alpha next adrenergic):ti,ab
#53.	(adrenergic next alpha):ti,ab
#54.	((alpha or alpha-adrenergic) near/2 block*).ti,ab
#55.	(or #14-#54)
#56.	#13 and #55

#### **B.1.2** Health Economics literature search strategy

- 2 Health economic evidence was identified by conducting a broad search relating to heart
- 3 valve disease population in NHS Economic Evaluation Database (NHS EED) (this ceased
- 4 to be updated after March 2015) and the Health Technology Assessment database (HTA) -
- 5 (this ceased to be updated after March 2018) with no date restrictions. NHS EED and HTA
- 6 databases are hosted by the Centre for Research and Dissemination (CRD). Additional
- 7 searches were run on Medline and Embase for health economics.

#### 8 Table 30: Database date parameters and filters used

Database	Dates searched	Search filter used
Medline	01 January 2014 – 15 October 2020	Exclusions Health economics studies
Embase	01 January 2014 – 15 October 2020	Exclusions Health economics studies
Centre for Research and Dissemination (CRD)	HTA - Inception – 31 March 2018 NHSEED - Inception to 31 March 2015	None

#### 9 Medline (Ovid) search terms

1.	exp Heart Valve Diseases/
2.	exp heart valves/
3.	((primary or secondary) adj valv* disease*).ti,ab.
4.	((valv* or flap* or leaflet*) adj1 (heart or cardiac) adj (disease* or disorder* or failure or failed or dysfunction* or insufficien* or repair* or replace* or damage* or leak*)).ti,ab.
5.	((mitral or aortic or tricuspid or pulmon*) adj (valv* or flap* or leaflet*) adj (disease* or disorder* or failure or failed or dysfunction* or insufficien* or repair* or replace* or damage* or leak*)).ti,ab.
6.	((mitral or aortic or tricuspid or pulmon*) adj3 (prolapse or regurgitation or stenos?s or atresia or insufficienc*)).ti,ab.
7.	Heart Valve Prosthesis/
8.	((mechanical or artificial or prosthe* or bioprosthe* or biological or tissue) adj (valv* or flap* or leaflet*)).ti,ab.
9.	valve-in-valve.ti,ab.
10.	(transcatheter adj2 (valve or valves)).ti,ab.

11.	exp Heart Murmurs/
12.	((heart or cardiac) adj murmur*).ti,ab.
13.	or/1-12
	letter/
14.	
15.	editorial/
16.	news/
17.	exp historical article/
18.	Anecdotes as Topic/
19.	comment/
20.	case report/
21.	(letter or comment*).ti.
22.	or/14-21
23.	randomized controlled trial/ or random*.ti,ab.
24.	22 not 23
25.	animals/ not humans/
26.	exp Animals, Laboratory/
27.	exp Animal Experimentation/
28.	exp Models, Animal/
29.	exp Rodentia/
30.	(rat or rats or mouse or mice).ti.
31.	or/24-30
32.	13 not 31
33.	limit 32 to English language
34.	(exp child/ or exp pediatrics/ or exp infant/) not (exp adolescent/ or exp adult/ or exp middle age/ or exp aged/)
35.	33 not 34
36.	Economics/
37.	Value of life/
38.	exp "Costs and Cost Analysis"/
39.	exp Economics, Hospital/
40.	exp Economics, Medical/
41.	Economics, Nursing/
42.	Economics, Pharmaceutical/
43.	exp "Fees and Charges"/
44.	exp Budgets/
45.	budget*.ti,ab.
46. 47.	cost*.ti.  (economic* or pharmaco?economic*).ti.
48.	(price* or pricing*).ti,ab.
48.	(cost* adj2 (effective* or utilit* or benefit* or minimi* or unit* or estimat* or
TJ.	variable*)).ab.
50.	(financ* or fee or fees).ti,ab.
51.	(value adj2 (money or monetary)).ti,ab.

52.	or/36-51
53.	35 and 52

# 1 Embase (Ovid) search terms

1.	exp valvular heart disease/
2.	exp heart valve/
3.	((primary or secondary) adj valv* disease*).ti,ab.
4.	((valv* or flap* or leaflet*) adj1 (heart or cardiac) adj (disease* or disorder* or failure or failed or dysfunction* or insufficien* or repair* or replace* or damage* or leak*)).ti,ab.
5.	((mitral or aortic or tricuspid or pulmon*) adj (valv* or flap* or leaflet*) adj (disease* or disorder* or failure or failed or dysfunction* or insufficien* or repair* or replace* or damage* or leak*)).ti,ab.
6.	((mitral or aortic or tricuspid or pulmon*) adj3 (prolapse or regurgitation or stenos?s or atresia or insufficienc*)).ti,ab.
7.	exp heart valve prosthesis/
8.	((mechanical or artificial or prosthe* or bioprosthe* or biological or tissue) adj (valv* or flap* or leaflet*)).ti,ab.
9.	valve-in-valve.ti,ab.
10.	(transcatheter adj2 (valve or valves)).ti,ab.
11.	exp heart murmur/
12.	((heart or cardiac) adj murmur*).ti,ab.
13.	or/1-12
14.	letter.pt. or letter/
15.	note.pt.
16.	editorial.pt.
17.	Case report/ or Case study/
18.	(letter or comment*).ti.
19.	or/14-18
20.	randomized controlled trial/ or random*.ti,ab.
21.	19 not 20
22.	animal/ not human/
23.	Nonhuman/
24.	exp Animal Experiment/
25.	exp Experimental animal/
26.	Animal model/
27.	exp Rodent/
28.	(rat or rats or mouse or mice).ti.
29.	or/21-28
30.	13 not 29
31.	limit 30 to English language
32.	(exp child/ or exp pediatrics/) not (exp adult/ or exp adolescent/)
33.	31 not 32
34.	health economics/
35.	exp economic evaluation/
36.	exp health care cost/
37.	exp fee/

38.	budget/
39.	funding/
40.	budget*.ti,ab.
41.	cost*.ti.
42.	(economic* or pharmaco?economic*).ti.
43.	(price* or pricing*).ti,ab.
44.	(cost* adj2 (effective* or utilit* or benefit* or minimi* or unit* or estimat* or variable*)).ab.
45.	(financ* or fee or fees).ti,ab.
46.	(value adj2 (money or monetary)).ti,ab.
47.	or/34-46
48.	33 and 47

#### 1 NHS EED and HTA (CRD) search terms

#1. MeSH DESCRIPTOR Heart Valve Diseases EXPLODE ALL TREES  #2. MeSH DESCRIPTOR Heart Valves EXPLODE ALL TREES  #3. (((primary or secondary) adj Valv* adj disease*))  #4. (((valv* or flap* or leaflet*) adj (heart or cardiac) adj (disease* or disorder* or failure or failed or dysfunction* or insufficien* or repair* or replace* or damage* or leak*)))  #5. ((heart or cardiac) adj (valv* or flap* or leaflet*) adj (disease* or disorder* or failure or failed or dysfunction* or insufficien* or repair* or replace* or damage* or leak*))  #6. (((mitral or aortic or tricuspid or pulmon*) adj (valv* or flap* or leaflet*) adj (disease* or disorder* or failure or failed or dysfunction* or insufficien* or repair* or replace* or damage* or leak*)))  #7. (((mitral or aortic or tricuspid or pulmon*) adj3 (prolapse or regurgitation or stenos?s or atresia or insufficienc*)))  #8. MeSH DESCRIPTOR Heart Valve Prosthesis EXPLODE ALL TREES  #9. (((mechanical or artificial or prosthe* or bioprosthe* or biological or tissue) adj (valv* or flap* or leaflet*)))  #10. (valve-in-valve)  #11. ((transcatheter adj2 (valve or valves)))	HO LED and HIA (OKD) Scaron terms	
#3. (((primary or secondary) adj Valv* adj disease*))  #4. (((valv* or flap* or leaflet*) adj (heart or cardiac) adj (disease* or disorder* or failure or failed or dysfunction* or insufficien* or repair* or replace* or damage* or leak*)))  #5. ((heart or cardiac) adj (valv* or flap* or leaflet*) adj (disease* or disorder* or failure or failed or dysfunction* or insufficien* or repair* or replace* or damage* or leak*))  #6. (((mitral or aortic or tricuspid or pulmon*) adj (valv* or flap* or leaflet*) adj (disease* or disorder* or failure or failed or dysfunction* or insufficien* or repair* or replace* or damage* or leak*)))  #7. (((mitral or aortic or tricuspid or pulmon*) adj3 (prolapse or regurgitation or stenos?s or atresia or insufficienc*)))  #8. MeSH DESCRIPTOR Heart Valve Prosthesis EXPLODE ALL TREES  #9. (((mechanical or artificial or prosthe* or bioprosthe* or biological or tissue) adj (valv* or flap* or leaflet*)))  #10. (valve-in-valve)	#1.	MeSH DESCRIPTOR Heart Valve Diseases EXPLODE ALL TREES
#4. (((valv* or flap* or leaflet*) adj (heart or cardiac) adj (disease* or disorder* or failure or failed or dysfunction* or insufficien* or repair* or replace* or damage* or leak*)))  #5. ((heart or cardiac) adj (valv* or flap* or leaflet*) adj (disease* or disorder* or failure or failed or dysfunction* or insufficien* or repair* or replace* or damage* or leak*))  #6. (((mitral or aortic or tricuspid or pulmon*) adj (valv* or flap* or leaflet*) adj (disease* or disorder* or failure or failed or dysfunction* or insufficien* or repair* or replace* or damage* or leak*)))  #7. (((mitral or aortic or tricuspid or pulmon*) adj3 (prolapse or regurgitation or stenos?s or atresia or insufficienc*)))  #8. MeSH DESCRIPTOR Heart Valve Prosthesis EXPLODE ALL TREES  #9. (((mechanical or artificial or prosthe* or bioprosthe* or biological or tissue) adj (valv* or flap* or leaflet*)))  #10. (valve-in-valve)	#2.	MeSH DESCRIPTOR Heart Valves EXPLODE ALL TREES
failed or dysfunction* or insufficien* or repair* or replace* or damage* or leak*)))  #5.	#3.	(((primary or secondary) adj Valv* adj disease*))
failed or dysfunction* or insufficien* or repair* or replace* or damage* or leak*))  #6.	#4.	
disorder* or failure or failed or dysfunction* or insufficien* or repair* or replace* or damage* or leak*)))  #7.	#5.	
atresia or insufficienc*)))  #8. MeSH DESCRIPTOR Heart Valve Prosthesis EXPLODE ALL TREES  #9. (((mechanical or artificial or prosthe* or biological or tissue) adj (valv* or flap* or leaflet*)))  #10. (valve-in-valve)	#6.	disorder* or failure or failed or dysfunction* or insufficien* or repair* or replace* or
#9. (((mechanical or artificial or prosthe* or bioprosthe* or biological or tissue) adj (valv* or flap* or leaflet*))) #10. (valve-in-valve)	#7.	
flap* or leaflet*))) #10. (valve-in-valve)	#8.	MeSH DESCRIPTOR Heart Valve Prosthesis EXPLODE ALL TREES
	#9.	, , , , , , , , , , , , , , , , , , , ,
#11. ((transcatheter adj2 (valve or valves)))	#10.	(valve-in-valve)
	#11.	((transcatheter adj2 (valve or valves)))
#12. #1 or #2 or #3 or #4 or #5 or #6 or #7 or #8 or #9 or #10 or #11	#12.	#1 or #2 or #3 or #4 or #5 or #6 or #7 or #8 or #9 or #10 or #11

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### **B.2** Valve disease with heart failure

- 5 <u>Heart valve disease search strategy 5- pharmacological management with heart failure</u>
- 6 This literature search strategy was used for the following reviews:
  - In adults with heart failure and concomitant heart valve disease, what is the clinical and cost effectiveness of ACE inhibitors, ARBs, beta blockers, calcium channel blockers, digoxin, diuretics and nitrates to improve clinical outcome?
- The literature searches for this review are detailed below and complied with the methodology outlined in Developing NICE guidelines: the manual. 112
- 12 For more information, please see the Methodology review published as part of the
- 13 accompanying documents for this guideline.

# B.2.11 Clinical search literature search strategy

- 2 Searches were constructed using a PICO framework where population (P) terms were
- 3 combined with Intervention (I) and in some cases Comparison (C) terms. Outcomes (O) are
- 4 rarely used in search strategies for interventions as these concepts may not be well
- 5 described in title, abstract or indexes and therefore difficult to retrieve. Search filters were
- 6 applied to the search where appropriate.

7

# 8 Table 31: Database date parameters and filters used

Database	Dates searched	Search filter used
Medline (OVID)	1946 - 14 October 2020	Exclusions Randomised controlled trials Systematic review studies
Embase (OVID)	1974 - 14 October 2020	Exclusions Randomised controlled trials Systematic review studies
The Cochrane Library (Wiley)	Cochrane Reviews to 2020 Issue 10 of 12 CENTRAL to 2020 Issue 10 of 12	None

#### 9 Medline (Ovid) search terms

94.	exp Heart Valve Diseases/
95.	exp heart valves/
96.	((primary or secondary) adj valv* disease*).ti,ab.
97.	((valv* or flap* or leaflet*) adj1 (heart or cardiac) adj (disease* or disorder* or failure or failed or dysfunction* or insufficien* or repair* or replace* or damage* or leak*)).ti,ab.
98.	((mitral or aortic or tricuspid or pulmon*) adj (valv* or flap* or leaflet*) adj (disease* or disorder* or failure or failed or dysfunction* or insufficien* or repair* or replace* or damage* or leak*)).ti,ab.
99.	((mitral or aortic or tricuspid or pulmon*) adj3 (prolapse or regurgitation or stenos?s or atresia or insufficienc*)).ti,ab.
100.	Heart Valve Prosthesis/
101.	((mechanical or artificial or prosthe* or bioprosthe* or biological or tissue) adj (valv* or flap* or leaflet*)).ti,ab.
102.	valve-in-valve.ti,ab.
103.	(transcatheter adj2 (valve or valves)).ti,ab.
104.	exp Heart Murmurs/
105.	((heart or cardiac) adj murmur*).ti,ab.
106.	or/1-12
107.	letter/
108.	editorial/
109.	news/
110.	exp historical article/
111.	Anecdotes as Topic/
112.	comment/
113.	case report/
114.	(letter or comment*).ti.
115.	or/14-21

<sup>&</sup>lt;Click this field on the first page and insert footer text if required>
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116.	randomized controlled trial/ or random*.ti,ab.	
117.	22 not 23	
118.	animals/ not humans/	
119.	exp Animals, Laboratory/	
120.	exp Animal Experimentation/	
121.	exp Models, Animal/	
122.	exp Rodentia/	
123.	(rat or rats or mouse or mice).ti.	
124.	or/24-30	
125.	13 not 31	
126.	limit 32 to English language	
127.	(exp child/ or exp pediatrics/ or exp infant/) not (exp adolescent/ or exp adult/ or exp middle age/ or exp aged/)	
128.	33 not 34	
129.	randomized controlled trial.pt.	
130.	controlled clinical trial.pt.	
131.	randomi#ed.ti,ab.	
132.	placebo.ab.	
133.	randomly.ti,ab.	
134.	Clinical Trials as topic.sh.	
135.	trial.ti.	
136.	or/36-42	
137.	Meta-Analysis/	
138.	exp Meta-Analysis as Topic/	
139.	(meta analy* or metanaly* or meta regression).ti,ab.	
140.	((systematic* or evidence*) adj3 (review* or overview*)).ti,ab.	
141.	(reference list* or bibliograph* or hand search* or manual search* or relevant journals).ab.	
142.	(search strategy or search criteria or systematic search or study selection or data extraction).ab.	
143.	(search* adj4 literature).ab.	
144.	(medline or pubmed or cochrane or embase or psychlit or psyclit or psychinfo or psycinfo or cinahl or science citation index or bids or cancerlit).ab.	
145.	cochrane.jw.	
146.	((multiple treatment* or indirect or mixed) adj2 comparison*).ti,ab.	
147.	or/44-52	
148.	35 and (43 or 54)	
149.	exp Angiotensin-Converting Enzyme Inhibitors/	
150.	(("angiotensin-converting enzyme" or ace) adj2 (inhibitor* or antagonist*)).ti,ab.	
151.	(Captopril or cilazapril or enalapril or enalaprilat or fosinopril or lisinopril or perindopril or quinapril or ramipril or teprotide or imidapril or trandolopril).ti,ab.	
152.	exp Angiotensin Receptor Antagonists/	
153.	(angiotensin adj3 receptor adj3 (antagonist* or blocker*)).ti,ab.	
154.	(1-Sarcosine-8-Isoleucine or amlodipine or irbesartan or olmesartan or saralasin or telmisartan or valsartan or azilsartan or candesartan or eprosartan or losartan).ti,ab.	
155.	exp Adrenergic beta-Antagonists/	

156.	(Acebutolol or Atenolol or Bisoprolol or Carvedilol or Celiprolol or Esmolol or Labetalol or Metoprolol or Nadolol or Nebivolol or Oxprenolol or Propranolol or Pindolol or Sotalol or Timolol).ti,ab.
157.	(beta adj3 block*).ti,ab.
158.	((beta-adrenoceptor or b-adrenoceptor or beta-adrenergic) adj (block* or antagonist*)).ti,ab.
159.	(b adj3 block*).ti,ab.
160.	(beta adj2 antagonist*).ti,ab.
161.	exp Calcium Channel Blockers/
162.	(calcium adj3 (block* or antagonis* or inhibit*)).ti,ab.
163.	calcium channel receptor block*.ti,ab.
164.	(amlodipine or amrinone or bencyclane or bepridil or carvedilol or cinnarizine or conotoxin* or diltiazem or felodipine or fendiline or flunarizine or gallopamil or isradipine or lamotrigine or mibefradil or nicardipine or nifedipine or nimodipine or nisoldipine or nitrendipine or perhexiline or pregabalin or prenylamine or risedronic acid or tiapamil or verapamil or zonisamide or atenolol or clevidipine or lacidipine or lercanidipine or dihyropyridine*).ti,ab.
165.	exp Digoxin/
166.	digoxin.ti,ab.
167.	exp Diuretics/
168.	((oral* or subcut* or IV or intravenous* or iv or infusion* or drip or drips or augment* or sequential* or loop or "high ceiling") adj6 diuretic*).ti,ab.
169.	(augment* adj diuresis).ti,ab.
170.	(acetazolamide or amiloride or bendroflumenethiazide or bumentanide or chlorothiazide or chlorothalidone or clopamide or cyclopenhiazide or ethacrynic acid or ethoxzolamide or furosemide or hydrochlorothiazide or hydroflumethiazide or indapamide or mefruside or methazolamide or methyclothiazide or metolazone or muzolimine or polythiazide or potassium citrate or spironolactone or ticrynafen or torsemide or triamterene or trichlormethiazide or xipamide or isosorbide or mannitol or co-amilozide or co-triamterzide or co-flumactone or eplerenone or co-amilofruse).ti,ab.
171.	exp Nitrates/
172.	Nitroglycerin/
173.	Nitroprusside/
174.	Isosorbide Dinitrate/
175.	(nitroglycerin or gtn or nitrate* or nitroprusside*).ti,ab.
176.	(glyceryl trinitrate or isosorbide dinitrate or isosorbide mononitrate).ti,ab.
177.	or/56-83
178.	55 and 84

#### 1 Embase (Ovid) search terms

96.	exp valvular heart disease/
97.	exp heart valve/
98.	((primary or secondary) adj valv* disease*).ti,ab.
99.	((valv* or flap* or leaflet*) adj1 (heart or cardiac) adj (disease* or disorder* or failure or failed or dysfunction* or insufficien* or repair* or replace* or damage* or leak*)).ti,ab.
100.	((mitral or aortic or tricuspid or pulmon*) adj (valv* or flap* or leaflet*) adj (disease* or disorder* or failure or failed or dysfunction* or insufficien* or repair* or replace* or damage* or leak*)).ti,ab.
101.	((mitral or aortic or tricuspid or pulmon*) adj3 (prolapse or regurgitation or stenos?s or atresia or insufficienc*)).ti,ab.
102.	exp heart valve prosthesis/

103.	((mechanical or artificial or prosthe* or bioprosthe* or biological or tissue) adj (valv* or flap* or leaflet*)).ti,ab.	
104.	valve-in-valve.ti,ab.	
105.	(transcatheter adj2 (valve or valves)).ti,ab.	
106.	exp heart murmur/	
107.	((heart or cardiac) adj murmur*).ti,ab.	
108.	or/1-12	
109.	letter.pt. or letter/	
110.	note.pt.	
111.	editorial.pt.	
112.	Case report/ or Case study/	
113.	(letter or comment*).ti.	
114.	or/14-18	
115.	randomized controlled trial/ or random*.ti,ab.	
116.	19 not 20	
117.	animal/ not human/	
118.	Nonhuman/	
119.	exp Animal Experiment/	
120.	exp Experimental animal/	
121.	Animal model/	
122.	exp Rodent/	
123.	(rat or rats or mouse or mice).ti.	
124.	or/21-28	
125.	13 not 29	
126.	limit 30 to English language	
127.	(exp child/ or exp pediatrics/) not (exp adult/ or exp adolescent/)	
128.	31 not 32	
129.	random*.ti,ab.	
130.	factorial*.ti,ab.	
131.	(crossover* or cross over*).ti,ab.	
132.	((doubl* or singl*) adj blind*).ti,ab.	
133.	(assign* or allocat* or volunteer* or placebo*).ti,ab.	
134.	crossover procedure/	
135.	single blind procedure/	
136.	randomized controlled trial/	
137.	double blind procedure/	
138.	or/34-42	
139.	systematic review/	
140.	meta-analysis/	
141.	(meta analy* or metanaly* or metaanaly* or meta regression).ti,ab.	
142.	((systematic or evidence) adj3 (review* or overview*)).ti,ab.	
143.	(reference list* or bibliograph* or hand search* or manual search* or relevant journals).ab.	
144.	(search strategy or search criteria or systematic search or study selection or data extraction).ab.	
145.	(search* adj4 literature).ab.	

146.	(medline or pubmed or cochrane or embase or psychlit or psyclit or psychinfo or psycinfo or cinahl or science citation index or bids or cancerlit).ab.	
147.	((pool* or combined) adj2 (data or trials or studies or results)).ab.	
148.	cochrane.jw.	
149.	((multiple treatment* or indirect or mixed) adj2 comparison*).ti,ab.	
150.	or/44-53	
151.	33 and (43 or 55)	
152.	exp dipeptidyl carboxypeptidase inhibitor/	
153.	(("angiotensin-converting enzyme" or ace) adj2 (inhibitor* or antagonist*)).ti,ab.	
154.	(Captopril or cilazapril or enalapril or enalaprilat or fosinopril or lisinopril or perindopril or quinapril or ramipril or teprotide or imidapril or trandolopril).ti,ab.	
155.	exp angiotensin receptor antagonist/	
156.	(angiotensin adj3 receptor adj3 (antagonist* or blocker*)).ti,ab.	
157.	(1-Sarcosine-8-Isoleucine or amlodipine or irbesartan or olmesartan or saralasin or telmisartan or valsartan or azilsartan or candesartan or eprosartan or losartan).ti,ab.	
158.	exp beta adrenergic receptor blocking agent/	
159.	(Acebutolol or Atenolol or Bisoprolol or Carvedilol or Celiprolol or Esmolol or Labetalol or Metoprolol or Nadolol or Nebivolol or Oxprenolol or Propranolol or Pindolol or Sotalol or Timolol).ti,ab.	
160.	(beta adj3 block*).ti,ab.	
161.	((beta-adrenoceptor or b-adrenoceptor or beta-adrenergic) adj (block* or antagonist*)).ti,ab.	
162.	(b adj3 block*).ti,ab.	
163.	(beta adj2 antagonist*).ti,ab.	
164.	exp Calcium Channel Blockers/	
165.	(calcium adj3 (block* or antagonis* or inhibit*)).ti,ab.	
166.	calcium channel receptor block*.ti,ab.	
167.	(amlodipine or amrinone or bencyclane or bepridil or carvedilol or cinnarizine or conotoxin* or diltiazem or felodipine or fendiline or flunarizine or gallopamil or isradipine or lamotrigine or mibefradil or nicardipine or nifedipine or nimodipine or nisoldipine or nitrendipine or perhexiline or pregabalin or prenylamine or risedronic acid or tiapamil or verapamil or zonisamide or atenolol or clevidipine or lacidipine or lercanidipine or dihyropyridine*).ti,ab.	
168.	digoxin/	
169.	digoxin.ti,ab.	
170.	exp diuretic agent/	
171.	((oral* or subcut* or IV or intravenous* or iv or infusion* or drip or drips or augment* or sequential* or loop or "high ceiling") adj6 diuretic*).ti,ab.	
172.	(augment* adj diuresis).ti,ab.	
173.	(acetazolamide or amiloride or bendroflumenethiazide or bumentanide or chlorothiazide or chlorothalidone or clopamide or cyclopenhiazide or ethacrynic acid or ethoxzolamide or furosemide or hydrochlorothiazide or hydroflumethiazide or indapamide or mefruside or methazolamide or methyclothiazide or metolazone or muzolimine or polythiazide or potassium citrate or spironolactone or ticrynafen or torsemide or triamterene or trichlormethiazide or xipamide or isosorbide or mannitol or co-amilozide or co-triamterzide or co-flumactone or eplerenone or co-amilofruse).ti,ab.	
174.	nitrate/	
175.	glyceryl trinitrate/	
176.	nitroprusside sodium/	
177.	isosorbide mononitrate/	

178.	isosorbide dinitrate/
179.	(nitroglycerin or gtn or nitrate* or nitroprusside*).ti,ab.
180.	(glyceryl trinitrate or isosorbide dinitrate or isosorbide mononitrate).ti,ab.
181.	or/57-85
182.	56 and 86

1 Cochrane Library (Wiley) search terms

#57.	MeSH descriptor: [Heart Valve Diseases] explode all trees
#58.	MeSH descriptor: [Heart Valves] explode all trees
#59.	((primary or secondary) NEXT valv* disease*):ti,ab
#60.	((valv* or flap* or leaflet*) near/2 (heart or cardiac) NEXT (disease* or disorder* or failure or failed or dysfunction* or insufficien* or repair* or replace* or damage* or leak*)):ti,ab
#61.	((mitral or aortic or tricuspid or pulmon*) NEXT (valv* or flap* or leaflet*) NEXT (disease* or disorder* or failure or failed or dysfunction* or insufficien* or repair* or replace* or damage* or leak*)):ti,ab
#62.	((mitral or aortic or tricuspid or pulmon*) NEAR/3 (prolapse or regurgitation or stenos?s or atresia or insufficienc*)):ti,ab
#63.	MeSH descriptor: [Heart Valve Prosthesis] explode all trees
#64.	((mechanical or artificial or prosthe* or bioprosthe* or biological or tissue) NEXT (valv* or flap* or leaflet*)):ti,ab
#65.	valve-in-valve:ti,ab
#66.	(transcatheter NEAR/2 (valve or valves)):ti,ab
#67.	MeSH descriptor: [Heart Murmurs] explode all trees
#68.	((heart or cardiac) NEXT murmur*):ti,ab
#69.	(or #1-#12)
#70.	MeSH descriptor: [Angiotensin-Converting Enzyme Inhibitors] explode all trees
#71.	(("angiotensin-converting enzyme" or ace) near/2 (inhibitor* or antagonist*)):ti,ab
#72.	(Captopril or cilazapril or enalapril or enalaprilat or fosinopril or lisinopril or perindopril or quinapril or ramipril or teprotide or imidapril or trandolopril):ti,ab
#73.	MeSH descriptor: [Angiotensin Receptor Antagonists] explode all trees
#74.	(angiotensin near/3 receptor near/3 (antagonist* or blocker*)):ti,ab
#75.	(amlodipine or irbesartan or olmesartan or saralasin or telmisartan or valsartan or azilsartan or candesartan or eprosartan or losartan):ti,ab
#76.	1-Sarcosine-8-Isoleucine:ti,ab
#77.	MeSH descriptor: [Adrenergic beta-Antagonists] explode all trees
#78.	(Acebutolol or Atenolol or Bisoprolol or Carvedilol or Celiprolol or Esmolol or Labetalol or Metoprolol or Nadolol or Nebivolol or Oxprenolol or Propranolol or Pindolol or Sotalol or Timolol):ti,ab
#79.	(beta near/3 block*):ti,ab
#80.	((beta-adrenoceptor or b-adrenoceptor or beta-adrenergic) NEXT (block* or antagonist*)):ti,ab
#81.	(b near/3 block*):ti,ab
#82.	(beta near/2 antagonist*):ti,ab
#83.	MeSH descriptor: [Calcium Channel Blockers] explode all trees
#84.	(calcium near/3 (block* or antagonis* or inhibit*)):ti,ab
#85.	'calcium channel receptor':ti,ab
#86.	(amlodipine or amrinone or bencyclane or bepridil or carvedilol or cinnarizine or conotoxin* or diltiazem or felodipine or fendiline or flunarizine or gallopamil or

	isradipine or lamotrigine or mibefradil or nicardipine or nifedipine or nimodipine or nisoldipine or nitrendipine or perhexiline or pregabalin or prenylamine or risedronic acid or tiapamil or verapamil or zonisamide or atenolol or clevidipine or lacidipine or lercanidipine or dihyropyridine*):ti,ab
#87.	MeSH descriptor: [Digoxin] explode all trees
#88.	digoxin:ti,ab
#89.	MeSH descriptor: [Diuretics] explode all trees
#90.	((oral* or subcut* or IV or intravenous* or iv or infusion* or drip or drips or augment* or sequential* or loop or "high ceiling") near/6 diuretic*):ti,ab
#91.	(acetazolamide or amiloride or bendroflumenethiazide or bumentanide or chlorothiazide or chlorothalidone or clopamide or cyclopenhiazide or ethacrynic acid or ethoxzolamide or furosemide or hydrochlorothiazide or hydroflumethiazide or indapamide or mefruside or methazolamide or methyclothiazide or metolazone or muzolimine or polythiazide or potassium citrate or spironolactone or ticrynafen or torsemide or triamterene or trichlormethiazide or xipamide or isosorbide or mannitol or co-amilozide or co-triamterzide or co-flumactone or eplerenone or co-amilofruse):ti,ab
#92.	MeSH descriptor: [Nitrates] explode all trees
#93.	MeSH descriptor: [Nitroglycerin] explode all trees
#94.	MeSH descriptor: [Nitroprusside] explode all trees
#95.	MeSH descriptor: [Isosorbide Dinitrate] explode all trees
#96.	(nitroglycerin or gtn or nitrate* or nitroprusside*):ti,ab
#97.	('glyceryl trinitrate' or 'isosorbide dinitrate' or 'isosorbide mononitrate'):ti,ab
#98.	(or #14-#41)
#99.	#13 and #42

### B.2.2 Health Economics literature search strategy

- 2 Health economic evidence was identified by conducting a broad search relating to heart
- 3 valve disease population in NHS Economic Evaluation Database (NHS EED) (this ceased
- 4 to be updated after March 2015) and the Health Technology Assessment database (HTA) -
- 5 (this ceased to be updated after March 2018) with no date restrictions. NHS EED and HTA
- 6 databases are hosted by the Centre for Research and Dissemination (CRD). Additional
- 7 searches were run on Medline and Embase for health economics.

#### 8 Table 32: Database date parameters and filters used

Database	Dates searched	Search filter used
Medline	01 January 2014 – 15 October 2020	Exclusions Health economics studies
Embase	01 January 2014 – 15 October 2020	Exclusions Health economics studies
Centre for Research and Dissemination (CRD)	HTA - Inception – 31 March 2018 NHSEED - Inception to 31 March 2015	None

#### 9 Medline (Ovid) search terms

54.	exp Heart Valve Diseases/	
55.	exp heart valves/	
56.	((primary or secondary) adj valv* disease*).ti,ab.	
57.	((valv* or flap* or leaflet*) adj1 (heart or cardiac) adj (disease* or disorder* or failure or failed or dysfunction* or insufficien* or repair* or replace* or damage* or leak*)).ti,ab.	

58.	((mitral or aortic or tricuspid or pulmon*) adj (valv* or flap* or leaflet*) adj (disease* or disorder* or failure or failed or dysfunction* or insufficien* or repair* or replace* or damage* or leak*)).ti,ab.	
59.	((mitral or aortic or tricuspid or pulmon*) adj3 (prolapse or regurgitation or stenos?s or atresia or insufficienc*)).ti,ab.	
60.	Heart Valve Prosthesis/	
61.	((mechanical or artificial or prosthe* or bioprosthe* or biological or tissue) adj (valv* or flap* or leaflet*)).ti,ab.	
62.	valve-in-valve.ti,ab.	
63.	(transcatheter adj2 (valve or valves)).ti,ab.	
64.	exp Heart Murmurs/	
65.	((heart or cardiac) adj murmur*).ti,ab.	
66.	or/1-12	
67.	letter/	
68.	editorial/	
69.	news/	
70.	exp historical article/	
71.	Anecdotes as Topic/	
72.	comment/	
73.	case report/	
74.	(letter or comment*).ti.	
75.	or/14-21	
76.	randomized controlled trial/ or random*.ti,ab.	
77.	22 not 23	
78.	animals/ not humans/	
79.	exp Animals, Laboratory/	
80.	exp Animal Experimentation/	
81.	exp Models, Animal/	
82.	exp Rodentia/	
83.	(rat or rats or mouse or mice).ti.	
84.	or/24-30	
	13 not 31	
85.		
86. 87.	limit 32 to English language  (exp child/ or exp pediatrics/ or exp infant/) not (exp adolescent/ or exp adult/ or exp middle age/ or exp aged/)	
88.	33 not 34	
89.	Economics/	
90.	Value of life/	
91.	exp "Costs and Cost Analysis"/	
92.	exp Economics, Hospital/	
93.	exp Economics, Medical/	
94.	Economics, Nursing/	
95.	Economics, Pharmaceutical/	
96.	exp "Fees and Charges"/	

97.	exp Budgets/
98.	budget*.ti,ab.
99.	cost*.ti.
100.	(economic* or pharmaco?economic*).ti.
101.	(price* or pricing*).ti,ab.
102.	(cost* adj2 (effective* or utilit* or benefit* or minimi* or unit* or estimat* or variable*)).ab.
103.	(financ* or fee or fees).ti,ab.
104.	(value adj2 (money or monetary)).ti,ab.
105.	or/36-51
106.	35 and 52

# 1 Embase (Ovid) search terms

49.	exp valvular heart disease/
50.	exp heart valve/
51.	((primary or secondary) adj valv* disease*).ti,ab.
52.	((valv* or flap* or leaflet*) adj1 (heart or cardiac) adj (disease* or disorder* or failure or failed or dysfunction* or insufficien* or repair* or replace* or damage* or leak*)).ti,ab.
53.	((mitral or aortic or tricuspid or pulmon*) adj (valv* or flap* or leaflet*) adj (disease* or disorder* or failure or failed or dysfunction* or insufficien* or repair* or replace* or damage* or leak*)).ti,ab.
54.	((mitral or aortic or tricuspid or pulmon*) adj3 (prolapse or regurgitation or stenos?s or atresia or insufficienc*)).ti,ab.
55.	exp heart valve prosthesis/
56.	((mechanical or artificial or prosthe* or bioprosthe* or biological or tissue) adj (valv* or flap* or leaflet*)).ti,ab.
57.	valve-in-valve.ti,ab.
58.	(transcatheter adj2 (valve or valves)).ti,ab.
59.	exp heart murmur/
60.	((heart or cardiac) adj murmur*).ti,ab.
61.	or/1-12
62.	letter.pt. or letter/
63.	note.pt.
64.	editorial.pt.
65.	Case report/ or Case study/
66.	(letter or comment*).ti.
67.	or/14-18
68.	randomized controlled trial/ or random*.ti,ab.
69.	19 not 20
70.	animal/ not human/
71.	Nonhuman/
72.	exp Animal Experiment/
73.	exp Experimental animal/
74.	Animal model/
75.	exp Rodent/
76.	(rat or rats or mouse or mice).ti.
77.	or/21-28

78.	13 not 29
79.	limit 30 to English language
80.	(exp child/ or exp pediatrics/) not (exp adult/ or exp adolescent/)
81.	31 not 32
82.	health economics/
83.	exp economic evaluation/
84.	exp health care cost/
85.	exp fee/
86.	budget/
87.	funding/
88.	budget*.ti,ab.
89.	cost*.ti.
90.	(economic* or pharmaco?economic*).ti.
91.	(price* or pricing*).ti,ab.
92.	(cost* adj2 (effective* or utilit* or benefit* or minimi* or unit* or estimat* or variable*)).ab.
93.	(financ* or fee or fees).ti,ab.
94.	(value adj2 (money or monetary)).ti,ab.
95.	or/34-46
96.	33 and 47

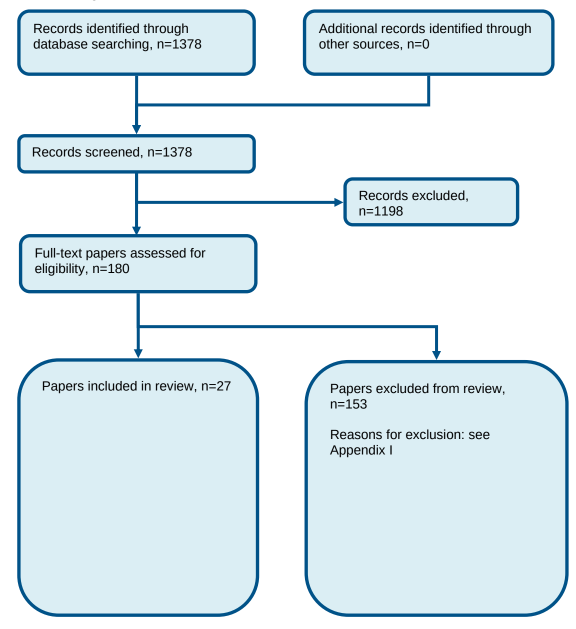
#### 1 NHS EED and HTA (CRD) search terms

#13.	MeSH DESCRIPTOR Heart Valve Diseases EXPLODE ALL TREES
#14.	MeSH DESCRIPTOR Heart Valves EXPLODE ALL TREES
#15.	((((primary or secondary) adj Valv* adj disease*))
#16.	(((valv* or flap* or leaflet*) adj (heart or cardiac) adj (disease* or disorder* or failure or failed or dysfunction* or insufficien* or repair* or replace* or damage* or leak*)))
#17.	((heart or cardiac) adj (valv* or flap* or leaflet*) adj (disease* or disorder* or failure or failed or dysfunction* or insufficien* or repair* or replace* or damage* or leak*))
#18.	(((mitral or aortic or tricuspid or pulmon*) adj (valv* or flap* or leaflet*) adj (disease* or disorder* or failure or failed or dysfunction* or insufficien* or repair* or replace* or damage* or leak*)))
#19.	(((mitral or aortic or tricuspid or pulmon*) adj3 (prolapse or regurgitation or stenos?s or atresia or insufficienc*)))
#20.	MeSH DESCRIPTOR Heart Valve Prosthesis EXPLODE ALL TREES
#21.	(((mechanical or artificial or prosthe* or bioprosthe* or biological or tissue) adj (valv* or flap* or leaflet*)))
#22.	(valve-in-valve)
#23.	((transcatheter adj2 (valve or valves)))
#24.	#1 or #2 or #3 or #4 or #5 or #6 or #7 or #8 or #9 or #10 or #11

# **Appendix C: Clinical evidence selection**

# **C.4** Valve disease without heart failure

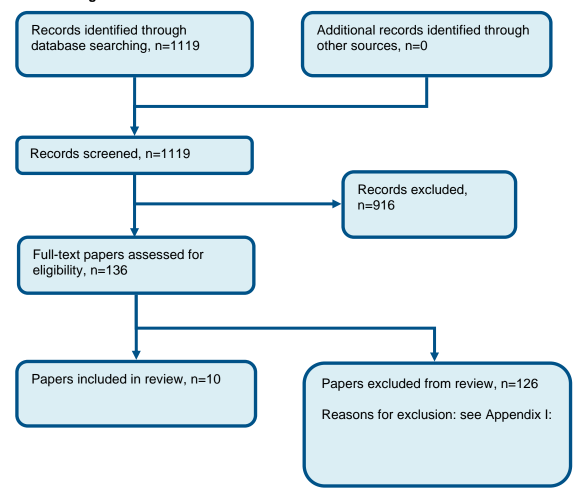
Figure 1: Flow chart of clinical study selection for the review of pharmacological management of heart valve disease without concomitant heart failure



1

# C.2 Valve disease with heart failure

Figure 2: Flow chart of clinical study selection for the review of pharmacological management



# **Appendix D: Clinical evidence tables**

Valve disease without heart failure

2

Study	Ahmed 2012 <sup>4</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=38)
Countries and setting	Conducted in USA; Setting: Outpatient follow up
Line of therapy	1st line
Duration of study	Intervention + follow up: 2 years
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: Echocardiography - colour flow Doppler imaging
Stratum	Primary mitral regurgitation
Subgroup analysis within study	Not applicable
Inclusion criteria	Moderate or severe MR documented by color flow Doppler, LVEF >55%, LV end systolic dimension (ESD) < 40 mm, and echocardiographic thickening of the mitral valve leaflets and prolapse
Exclusion criteria	New York Heart Association class III or IV symptoms, previous myocardial infarction, significant coronary artery disease by exercise testing with myocardial perfusion imaging, significant other valvular disease, serum creatinine >2.5, and hypertension requiring medical treatment
Recruitment/selection of patients	Recruited in Birmingham, Alabama. No other details given.
Age, gender and ethnicity	Age - Mean (SD): Metoprolol: 52.9 (9.1). Placebo: 56 (9.2). Gender (M:F): 18:20. Ethnicity: Majority (92%) Caucasian
Further population details	1. Age: <75 years (Metoprolol: 52.9 (9.1). Placebo: 56 (9.2).). 2. Disease mechanism for aortic and mitral stenosis: Not applicable 3. Presence vs. absence of uncontrolled systemic hypertension (140/85 mmHg) at the end of trial intervention despite medication: Not stated / Unclear (At baseline: Metoprolol: 125/75 (14/8). Placebo: 121/75 (14/11).).
Indirectness of population	No indirectness
Interventions	(n=19) Intervention 1: Beta blockers - Metoprolol . Toprol XL (range 25 to 100mg/day) - starting dose of 12.5-25mg/day titrated up to the maximum tolerable dose at 2 week intervals up to a maximum of 100mg/day. Duration 2 years. Concurrent medication/care: No additional information given. Indirectness: No indirectness

Heart valve disease: DRAFT FOR CONSULTATION 4 MIDs used to assess imprecision were ±4.0

	(n=19) Intervention 2: Placebo. Placebo. Duration 2 years. Concurrent medication/care: No additional information given. Indirectness: No indirectness
Funding	Equipment / drugs provided by industry (Drug and placebo supplied by Astra- Zenica. The study was funded by NHLBI Specialised Centre for Clinically Oriented Research (SCCOR) in Cardiac Dysfunction)

4 MIDs used to assess imprecision were ±4.0 disease: DRAFT

FO.

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#### RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: METOPROLOL versus PLACEBO

Protocol outcome 1: All-cause mortality at ≥12 months

- Actual outcome for Primary mitral regurgitation: Death at 2 years; Group 1: 1/19, Group 2: 0/18; Comments: 1 death due to PE after a cosmetic procedure

Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 0; Group 2 Number missing: 1, Reason: 1 person withdrew from the study early on

Protocol outcome 2: Cardiac mortality at ≥12 months

- Actual outcome for Primary mitral regurgitation: Death at 2 years; Group 1: 1/19, Group 2: 0/18; Comments: 1 death due to PE after a cosmetic procedure

Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 0; Group 2 Number missing: 1, Reason: 1 person withdrew from the study early on

Protocol outcome 3: Need for heart valve intervention at ≥12 months

- Actual outcome for Primary mitral regurgitation: Need to have valve replacement or surgery at 2 years; Group 1: 2/18, Group 2: 6/18 Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 1, Reason: 1 death; Group 2 Number missing: 1, Reason: 1 person withdrew from the study early on

Protocol outcome 4: Withdrawal due to adverse events at ≥12 months

- Actual outcome for Primary mitral regurgitation: Serious adverse events at 2 years; Group 1: 3/18, Group 2: 7/18; Comments: Events not stated Risk of bias: All domain - Very high, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: Serious indirectness, Comments: Doesn't report withdrawal due to adverse events; Group 1 Number missing: 1, Reason: 1 death; Group 2 Number missing: 1, Reason: 1 person withdrew from the study early on

Protocol outcomes not reported by the study	Quality of life at 6 months; Quality of life at ≥12 months; Onset of symptoms or progression of NYHA class at ≥12 months; Evidence of HVD progression on imaging (worsening of disease severity) at ≥12 months; Exercise tolerance at ≥12 months; Withdrawal due to adverse events at 6 months
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Study (subsidiary papers)	ASTRONOMER trial: Chan 2010 <sup>27</sup> (Chan 2011 <sup>25</sup> , Chan 2010 <sup>26</sup> , Chan 2007 <sup>28</sup> )	
Study type	RCT (Patient randomised; Parallel)	
Number of studies (number of participants)	1 (n=272)	
Countries and setting	Conducted in Canada; Setting: Outpatient follow up in secondary care	
ine of therapy	1st line	
Duration of study	Intervention + follow up: Median follow-up: 3.5 years (IQR: 2.1 to 4.5 years)	
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: Echocardiography - people were recruited from echocardiographic laboratories and cardiology clinics at participating sites.	
Stratum	Primary aortic [including bicuspid] stenosis: While stated as including mild to moderate aortic stenosis, the mean peak aortic valve velocity was 3.16 (0.42) in the intervention arm, and 3.19 (0.42) in the control arm, considered moderate severity by the British Society of Echocardiography guidelines.	
Subgroup analysis within study	Not applicable:	
nclusion criteria	People between 18 and 82 years of age with asymptomatic mild to moderate aortic stenosis defined by maximum aortic valve velocity between 2.5 and 4.0m/s.	
Exclusion criteria	People with clinical indications for the use of statin as defined by Canadian guidelines such as coronary artery disease, cerebrovascular disease, peripheral vascular disease and diabetes. Baseline lipid values outside of the target levels for their respective risk category according to Canadian guidelines.	
Recruitment/selection of patients	People recruited from echocardiographic laboratories and cardiology clinics at participating sites (5 sites listed in the study).	
Age, gender and ethnicity	Age - Mean (SD): 57.9 (13.6). Gender (M:F): 166:103. Ethnicity: 98% were white. Asian people were excluded for the later part of the study due to concern regarding adverse events. 1 Asian person remained in the study due to no adverse events.	
Further population details	1. Age: <75 years 2. Disease mechanism for aortic and mitral stenosis: Mixed (Not completely stated. However, they do state that around 50% of people had bicuspid aortic valve disease.). 3. Presence vs. absence of uncontrolled systemic hypertension (140/85 mmHg) at the end of trial intervention despite medication: Not stated / Unclear (Mean blood pressure at the start of study was not hypertensive. Intervention: 128.8/76.5 (15.67/10.04). Control: 128.4/75.9 (15.94/10.92).).	
ndirectness of population	No indirectness	

Interventions	(n=136) Intervention 1: Statins - Rosuvastatin . Rosuvastatin 40mg daily. Duration 3.5 years. Concurrent medication/care: No information provided. Indirectness: No indirectness  (n=136) Intervention 2: Placebo. Placebo. Duration 3.5 years. Concurrent medication/care: No information provided. Indirectness: No indirectness
Funding	Study funded by industry (Supported by the Canada Institutes of Health Research, with additional support from AstraZeneca Canada Inc. Also several authors received travel grants from Astra Zeneca. One author has had consultancies for and/or research funds from St Jude Medical, Edward Life Sciences, and Medtronic.)

#### RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: ROSUVASTATIN versus PLACEBO

Protocol outcome 1: All-cause mortality at ≥12 months

- Actual outcome for Primary aortic [including bicuspid] stenosis: All-cause mortality at 3.5 years; Group 1: 3/134, Group 2: 5/135
Risk of bias: All domain - Low, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 2, Reason: 136 allocated to rosuvastatin. 2 had a randomisation error and weren't eligible for randomisation - making 134 people who received rosuvastatin (included in intention to treat analysis). However, in actuality 57 people discontinued treatment: 25 for adverse events, 8 withdrew consent, 1 lost to follow-up, 1 investigator discretion, 3 patients died and 19 other (unclear).; Group 2 Number missing: 1, Reason: 136 allocated to placebo. 1 had a randomisation error and weren't eligible for randomisation - making 135 people who received placebo (included in the intention to treat analysis). However, in actuality 66 people discontinued treatment. 26 for adverse events. 2 for protocol non-compliance. 4 withdrew consent. 2 at investigator discretion. 5 patients died. 25 other (unclear).

Protocol outcome 2: Cardiac mortality at ≥12 months

- Actual outcome for Primary aortic [including bicuspid] stenosis: Cardiac death at 3.5 years; Group 1: 2/134, Group 2: 5/135
  Risk of bias: All domain Low, Selection Low, Blinding Low, Incomplete outcome data Low, Outcome reporting Low, Measurement Low, Crossover Low, Subgroups Low, Other 1 Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 2, Reason: 136 allocated to rosuvastatin. 2 had a randomisation error and weren't eligible for randomisation making 134 people who received rosuvastatin (included in intention to treat analysis). However, in actuality 57 people discontinued treatment: 25 for adverse events, 8 withdrew consent, 1 lost to follow-up, 1 investigator discretion, 3 patients died and 19 other (unclear).
- ; Group 2 Number missing: 1, Reason: 136 allocated to placebo. 1 had a randomisation error and weren't eligible for randomisation making 135 people who received placebo (included in the intention to treat analysis). However, in actuality 66 people discontinued treatment. 26 for adverse events. 2 for protocol non-compliance. 4 withdrew consent. 2 at investigator discretion. 5 patients died. 25 other (unclear).

Protocol outcome 3: Need for heart valve intervention at ≥12 months

- Actual outcome for Primary aortic [including bicuspid] stenosis: Aortic valve replacement at 3.5 years; Group 1: 28/134, Group 2: 27/135
  Risk of bias: All domain Low, Selection Low, Blinding Low, Incomplete outcome data Low, Outcome reporting Low, Measurement Low, Crossover Low, Subgroups Low, Other 1 Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 2, Reason: 136 allocated to rosuvastatin. 2 had a randomisation error and weren't eligible for
- randomisation making 134 people who received rosuvastatin (included in

intention to treat analysis). However, in actuality 57 people discontinued treatment: 25 for adverse events, 8 withdrew consent, 1 lost to follow-up, 1 investigator discretion, 3 patients died and 19 other (unclear).; Group 2 Number missing: 1, Reason: 136 allocated to placebo. 1 had a randomisation error and weren't eligible for randomisation - making 135 people who received placebo (included in the intention to treat analysis). However, in actuality 66 people discontinued

treatment. 26 for adverse events. 2 for protocol non-compliance. 4 withdrew consent. 2 at investigator discretion. 5 patients died. 25 other (unclear).

Protocol outcome 4: Withdrawal due to adverse events at ≥12 months

- Actual outcome for Primary aortic [including bicuspid] stenosis: Withdrawal due to adverse events at 3.5 years; Group 1: 25/134, Group 2: 26/135
  Risk of bias: All domain Low, Selection Low, Blinding Low, Incomplete outcome data Low, Outcome reporting Low, Measurement Low, Crossover
- Low, Subgroups Low, Other 1 Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 2, Reason: 136 allocated to rosuvastatin. 2 had a randomisation error and weren't eligible for

randomisation - making 134 people who received rosuvastatin (included in

intention to treat analysis). However, in actuality 57 people discontinued treatment: 25 for adverse events, 8 withdrew consent, 1 lost to follow-up, 1 investigator discretion, 3 patients died and 19 other (unclear).; Group 2 Number missing: 1, Reason: 136 allocated to placebo. 1 had a randomisation error and weren't eligible for randomisation - making 135 people who received placebo (included in the intention to treat analysis). However, in actuality 66 people discontinued

treatment. 26 for adverse events. 2 for protocol non-compliance. 4 withdrew consent. 2 at investigator discretion. 5 patients died. 25 other (unclear).

Protocol outcomes not reported by the study

Quality of life at 6 months; Quality of life at ≥12 months; Onset of symptoms or progression of NYHA class at ≥12 months; Evidence of HVD progression on imaging (worsening of disease severity) at ≥12 months; Exercise tolerance at ≥12 months; Withdrawal due to adverse events at 6 months

Study	Banaszewski 1998 <sup>13</sup>	
Study type	RCT (Patient randomised; Parallel)	
Number of studies (number of participants)	1 (n=31)	
Countries and setting	Conducted in Poland; Setting: Initially secondary care, followed by outpatient follow up	
Line of therapy	1st line	
Duration of study	Intervention + follow up: 2.75 years	
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: Echocardiography and cardiac catheterisation performed during the study - used echocardiographic parameters to determine severity of aortic regurgitation and exclude presence of other valve disease	
Stratum	Primary aortic regurgitation	
Subgroup analysis within study	Not applicable:	
Inclusion criteria	Known history of aortic regurgitation for >24 months with a stable clinical course during that time; no clinical symptoms of heart failure, and echo-Doppler parameters of isolated (at least moderate) aortic insufficiency, LVEDD >56mm and LVEF >50%; signed, written informed consent	
Exclusion criteria	A maximum aortic valvular pressure gradient >15mmHg; insufficiency and/or stenosis of any other valve; sustained or paroxysmal supra- and/or ventricular arrhythmias; coexistent coronary artery disease; previous therapy with ACE inhibitors, calcium channel blockers, diuretics, beta-blockers and/or digitalis	
Recruitment/selection of patients	Does not give any additional information about where people were recruited from	
Age, gender and ethnicity	Age - Mean (SD): 34.9 (10.1) (range: 18-60). Gender (M:F): 27:4. Ethnicity: Not stated	
Further population details	1. Age: <75 years (Mean age: 34.9 (10.1) (range: 18-60)). 2. Disease mechanism for aortic and mitral stenosis: Not applicable 3. Presence vs. absence of uncontrolled systemic hypertension (140/85 mmHg) at the end of trial intervention despite medication: Not stated / Unclear (At the start of the trial: Mean systolic blood pressure 132.2 (14.3). Mean diastolic blood pressure 54.5 (10.4). Does not report at the end.).	
Indirectness of population	No indirectness	
Interventions	(n=12) Intervention 1: Calcium-channel blockers (CCB) - Nifedipine. Nifedipine 10-20mg three times a day - mean daily dose of 40mg. People were given the	

Heart valve disease: DRAFT FOR CONSULTATION 4 MIDs used to assess imprecision were ±4.0

	maximum tolerable dose. Duration 2.75 years. Concurrent medication/care: Exclusion criteria stated that people should not have been using ACE inhibitors, calcium channel blockers, diuretics, beta-blockers and/or digitalis prior to the study. No other information available. Indirectness: No indirectness  (n=13) Intervention 2: Angiotensin-converting enzyme (ACE) inhibitors - Captopril. Captopril 12.5-30mg three times a day - mean daily dose of 75mg. People were given the maximum tolerable dose. Duration 2.75 years. Concurrent medication/care: Exclusion criteria stated that people should not have been using ACE inhibitors, calcium channel blockers, diuretics, beta-blockers and/or digitalis prior to the study. No other information available. Indirectness: No indirectness Comments: The study included an acute phase where participants underwent cardiac catheterisation and exercise therapy after a single dose of nifedipine, which was repeated after 24 hours (6 half-lives) with captopril. Another 24 hours were given and then the long term (randomised) phase of the study was started.
Funding	Funding not stated

RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: NIFEDIPINE versus CAPTOPRIL

Protocol outcome 1: Onset of symptoms or progression of NYHA class at ≥12 months

- Actual outcome for Primary aortic regurgitation: Development of symptoms at 2.75 years; Group 1: 0/12, Group 2: 0/13
Risk of bias: All domain - Very high, Selection - Very high, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low,

Risk of bias: All domain - Very high, Selection - Very high, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: Reports the baseline characteristics for the combined cohort, but not for each intervention group.; Group 1 Number missing: 0, Reason: 5 people withdrew consent after the acute phase and 1 withdrew due to adverse events during the acute phase. However, it is not reported whether there was randomisation before this withdrew due to adverse events during the acute phase. However, it is not reported whether there was randomisation before this withdrawal so assuming no missing people.

Protocol outcome 2: Evidence of HVD progression on imaging (worsening of disease severity) at ≥12 months

- Actual outcome for Primary aortic regurgitation: Aortic regurgitation grade worsening (by ≥1) at 2.75 years; Group 1: 2/12, Group 2: 0/13 Risk of bias: All domain - Very high, Selection - Very high, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: Reports the baseline characteristics for the combined cohort, but not for each intervention group.; Group 1 Number missing: 0, Reason: 5 people withdrew consent after the acute phase and 1 withdrew due to adverse events during the acute phase. However, it is not reported whether there was randomisation before this withdrew due to adverse events during the acute phase. However, it is not reported whether there was randomisation before this withdrawal so assuming no missing people.

Protocol outcomes not reported by the study	All-cause mortality at ≥12 months; Cardiac mortality at ≥12 months; Quality of life at 6 months; Quality of life at ≥12 months; Need for heart valve intervention at ≥12
	months; Exercise tolerance at ≥12 months; Withdrawal due to adverse events at 6 months; Withdrawal due to adverse events at ≥12 months

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Study	Broch 2016 <sup>20</sup>	
Study type	RCT (Patient randomised; Parallel)	
Number of studies (number of participants)	1 (n=75)	
Countries and setting	Conducted in Denmark, Norway; Setting: Outpatient follow up	
Line of therapy	1st line	
Duration of study	Intervention + follow up: 6 months	
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: Cardiac MRI and echocardiography	
Stratum	Primary aortic regurgitation	
Subgroup analysis within study	Not applicable	
Inclusion criteria	People aged between 18 and 70 years with asymptomatic, haemodynamically significant aortic regurgitation, an LVEF >50% and an LVEDD >5.0cm (or an indexed value >30cm/m²).	
Exclusion criteria	Symptoms of heart failure; a history of myocardial infarction or symptomatic coronary heart disease; significant aortic stenosis (valvular area <1.5cm²); additional haemodynamically significant valvular or congenital heart disease; an indication for aortic valve surgery (severe AR in conjunction with either symptoms of heart failure, an LVEF <50%, or an LV end diastolic/end systolic internal diameter >7.0/5.0cm); a second or third degree atrioventricular block; atrial fibrillation; an intra-cardiac device; serum creatinine >250 micromol/L; alanine aminotransferase or aspartate aminotransferase >3 times the upper limit of normal; any illness of disorder that could severely limit survival; conditions or circumstances likely to lead to poor treatment adherence; and intolerance to metoprolol CR/XL.	
Recruitment/selection of patients	Recruited from two centers.	
Age, gender and ethnicity	Age - Mean (SD): 44 (14). Gender (M:F): 67:8. Ethnicity: Not stated	
Further population details	1. Age: <75 years (44 (14)). 2. Disease mechanism for aortic and mitral stenosis: Not applicable 3. Presence vs. absence of uncontrolled systemic hypertension (140/85 mmHg) at the end of trial intervention despite medication: Absence of uncontrolled systemic hypertension (Final metoprolol: 124/58 (17/9). Final placebo: 134/67 (19/6).).	
Extra comments	55 (73%) people had bicuspid aortic valves	
Indirectness of population	No indirectness	
Interventions	(n=37) Intervention 1: Beta blockers - Metoprolol . Metoprolol CR/XL 25mg doubled every week up to a target daily dose of 200mg or the maximum tolerable dose.	

	Duration 6 months. Concurrent medication/care: All people were allowed to concomitantly use other vasoactive drugs. 6 on and ACE inhibitor/ARB. 2 on calcium channel blockers. 5 on statins. 7 on acetylsalicylic acid. 1 on another cardiovascular drug. Indirectness: No indirectness  (n=38) Intervention 2: Placebo. Placebo. Duration 6 months. Concurrent medication/care: All people were allowed to concomitantly use other vasoactive drugs. 6 on and ACE inhibitor/ARB. 3 on calcium channel blockers. 5 on statins. 2 on acetylsalicylic acid. Indirectness: No indirectness
Funding	Equipment / drugs provided by industry (Active drug and placebo provided by AstraZeneca. Unrestricted grants provided by the South-East Norway regional health authority and the Norwegian ExtraFoundation for Health and Rehabilitation through EXTRA funds.)

#### RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: METOPROLOL versus PLACEBO

Number missing: 2, Reason: 1 withdrew, 1 had a poor quality baseline MRI so their results were excluded

Protocol outcome 1: Quality of life at 6 months

- Actual outcome for Primary aortic regurgitation: EuroQol VAS (0-100) at 6 months; Group 1: mean 85 (SD 7); n=36, Group 2: mean 82 (SD 16); n=36; EuroQoL visual analogue scale 0-100 Top=High is good outcome; Comments: Baseline metoprolol: 84 (9). Baseline placebo: 82 (11). Risk of bias: All domain High, Selection High, Blinding Low, Incomplete outcome data Low, Outcome reporting Low, Measurement Low, Crossover Low, Subgroups Low, Other 1 Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 1, Reason: 1 withdrew; Group 2
- Actual outcome for Primary aortic regurgitation: KCCQ overall clinical summary score at 6 months; Group 1: mean 98 (SD 42.78); n=36, Group 2: mean 96 (SD 42.78); n=36; KCCQ 0-100 Top=High is good outcome; Comments: SD calculated from p value. Reported p-value = 0.78. Standard error = 7.13. Risk of bias: All domain High, Selection High, Blinding Low, Incomplete outcome data Low, Outcome reporting Low, Measurement Low, Crossover Low, Subgroups Low, Other 1 Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 1, Reason: 1 withdrew; Group 2 Number missing: 2, Reason: 1 withdrew, 1 had a poor quality baseline MRI so their results were excluded

#### Protocol outcome 2: Exercise tolerance at ≥12 months

- Actual outcome for Primary aortic regurgitation : Peak work (Watts) at 6 months; Group 1: mean 229 Watts (SD 62); n=36, Group 2: mean 241 Watts (SD 62); n=36; Comments: Baseline metoprolol: 240 (63). Baseline placebo: 237 (63).

Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 1, Reason: 1 withdrew; Group 2 Number missing: 2, Reason: 1 withdrew, 1 had a poor quality baseline MRI so their results were excluded

Protocol outc	omes not repor	rted by the study
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All-cause mortality at ≥12 months; Cardiac mortality at ≥12 months; Quality of life at ≥12 months; Onset of symptoms or progression of NYHA class at ≥12 months; Evidence of HVD progression on imaging (worsening of disease severity) at ≥12 months; Need for heart valve intervention at ≥12 months; Withdrawal due to adverse events at ≥12 months

Study	Bull 2015 <sup>21</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=100)
Countries and setting	Conducted in United Kingdom; Setting: Outpatient follow up
Line of therapy	1st line
Duration of study	Intervention + follow up: 1 year
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: Cardiovascular magnetic resonance imaging and echocardiography
Stratum	Primary aortic [including bicuspid] stenosis
Subgroup analysis within study	Not applicable
Inclusion criteria	All people aged >18 years with moderate or severe aortic stenosis by standard echocardiographic criteria (valve area <1.5cm², or peak velocity >3.0m/s (peak valve gradient >36mmHg)), who were asymptomatic as judged by patient-reported symptoms, and who did not have indications for valve replacement surgery.
Exclusion criteria	Abnormal LV function (LVEF <50% by echocardiography), other significant (>mild) valvular heart disease, excess hypo- or hypertension, intolerance to ACE inhibitors or ARBs or their prescription over the previous 3 months.
Recruitment/selection of patients	People recruited from clinics at the John Radcliffe Hospital and surrounding institutions.
Age, gender and ethnicity	Age - Mean (SD): 68.57 (14.22) years. Gender (M:F): 71:25. Ethnicity: Not stated
Further population details	1. Age: Mixed (68.57 (14.22) years. Confidence intervals would fall over the age boundary.). 2. Disease mechanism for aortic and mitral stenosis: Not stated / Unclear 3. Presence vs. absence of uncontrolled systemic hypertension (140/85 mmHg) at the end of trial intervention despite medication: Not stated / Unclear (At the start of the study 11 people in the ramipril arm and 17 people in the placebo arm had hypertension. No statement about the degree or progression during the study.).
Indirectness of population	No indirectness
Interventions	(n=50) Intervention 1: Angiotensin-converting enzyme (ACE) inhibitors - Ramipril. Ramipril 2.5mg daily for 2 weeks, raised to 5mg daily until the 3-month check, raised to 10mg daily for the rest of the study or to the maximal dose with no adverse events. Duration 1 year. Concurrent medication/care: Not stated. Indirectness: No indirectness

	(n=50) Intervention 2: Placebo. Placebo. Duration 1 year. Concurrent medication/care: Not stated. Indirectness: No indirectness
Funding	Academic or government funding (Funded by a grant from Heart Research UK and supported by the Oxford Comprehensive Biomedical Research Centre, funded by the UK National Institute for Health Research. One author was supported by a British Heart Foundation Clinical Research Training Fellowship.)

### RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: RAMIPRIL versus PLACEBO

Protocol outcome 1: Need for heart valve intervention at ≥12 months

- Actual outcome for Primary aortic [including bicuspid] stenosis: Aortic valve replacement at 12 months; Group 1: 4/40, Group 2: 2/43 Risk of bias: All domain - Very high, Selection - Very high, Blinding - Low, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: Placebo group generally has more hypertension, more medication usage and a worse exercise distance at baseline than the ramipril group.; Group 1 Number missing: 10, Reason: 50 allocated to ramipril. 1 withdrew consent before receiving the intervention. After receiving the allocated intervention: 6 withdrew consent, 2 withdrew due to a cough, 4 had an aortic valve replacement, 1 had a pacemaker implanted.; Group 2 Number missing: 7, Reason: 50 allocated to placebo. 2 withdrew consent and 1 had claustrophobia so withdrew from the study before receiving the intervention. After receiving the allocated intervention: 1 withdrew from the trial due to instruction from treating clinician, 2 withdrew consent, 1 withdrew due to a serious adverse event, 2 had an aortic valve replacement.

#### Protocol outcome 2: Exercise tolerance at ≥12 months

- Actual outcome for Primary aortic [including bicuspid] stenosis: Exercise distance (m) - measured with treadmill exercise test at 12 months; Group 1: mean -20 m (SD 26); n=26, Group 2: mean 29 m (SD 25); n=41; Comments: Baseline values ramipril: 1030 (386)m. Baseline values control: 985 (360)m. Risk of bias: All domain - Very high, Selection - Very high, Blinding - Low, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: Placebo group generally has more hypertension, more medication usage and a worse exercise distance at baseline than the ramipril group.; Group 1 Number missing: 14, Reason: 50 allocated to ramipril. 1 withdrew consent before receiving the intervention. After receiving the allocated intervention: 6 withdrew consent, 2 withdrew due to a cough, 4 had an aortic valve replacement, 1 had a pacemaker implanted.; Group 2 Number missing: 9, Reason: 50 allocated to placebo. 2 withdrew consent and 1 had claustrophobia so withdrew from the study before receiving the intervention. After receiving the allocated intervention: 1 withdrew from the trial due to instruction from treating clinician, 2 withdrew consent, 1 withdrew due to a serious adverse event, 2 had an aortic valve replacement.

### Protocol outcome 3: Withdrawal due to adverse events at ≥12 months

- Actual outcome for Primary aortic [including bicuspid] stenosis: Withdrawal due to adverse events at 12 months; Group 1: 2/38, Group 2: 1/42 Risk of bias: All domain - Very high, Selection - Very high, Blinding - Low, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: Placebo group generally has more hypertension, more medication usage and a worse exercise distance at baseline than the ramipril group.; Group 1 Number missing: 12, Reason: 50 allocated to ramipril. 1 withdrew consent before receiving the intervention. After receiving the allocated intervention: 6 withdrew consent, 2 withdrew due to

consent and 1 had claustrophobia so withdrew from the study before	planted.; Group 2 Number missing: 8, Reason: 50 allocated to placebo. 2 withdrew e receiving the intervention. After receiving the allocated intervention: 1 withdrew from t, 1 withdrew due to a serious adverse event, 2 had an aortic valve replacement.
Protocol outcomes not reported by the study	All-cause mortality at ≥12 months; Cardiac mortality at ≥12 months; Quality of life at ≥12 months; Quality of life at 6 months; Onset of symptoms or progression of NYHA class at ≥12 months; Evidence of HVD progression on imaging (worsening of disease severity) at ≥12 months; Withdrawal due to adverse events at 6 months

Study	Evangelista 2005 <sup>49</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=95)
Countries and setting	Conducted in Spain; Setting: Outpatient follow up
Line of therapy	1st line
Duration of study	Intervention + follow up: 7 years (range: 0.6 to 8.8 years)
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: Physical examination, echocardiography, 12-lead ECG, chest radiography and radionuclide angiography at rest.
Stratum	Primary aortic regurgitation
Subgroup analysis within study	Not applicable
Inclusion criteria	People with asymptomatic, chronic, severe aortic regurgitation (jet width exceeding 10mm and apical jet area exceeding 7cm <sup>2</sup> on colour Doppler ultrasonography, or when regurgitant fraction >60%) and normal left ventricular function.
Exclusion criteria	A decreased LVEF (<50%) during the preceding 6 months, other clinically significant associated valvular disease, associated valvular aortic stenosis (aortic mean gradient, more than 20mmHg), a diastolic blood pressure of more than 90mmHg, atrial fibrillation, or a history of coronary heart disease or other associated diseases that could affect the prognosis or functional class (including Marfan's syndrome or an ascending aortic aneurysm).
Recruitment/selection of patients	Consecutive people seen in the outpatient clinic
Age, gender and ethnicity	Age - Mean (SD): 44.35 (13.19). Gender (M:F): 74:21. Ethnicity: Not stated
Further population details	1. Age: <75 years 2. Disease mechanism for aortic and mitral stenosis: Not applicable 3. Presence vs. absence of uncontrolled systemic hypertension (140/85 mmHg) at the end of trial intervention despite medication: Presence of uncontrolled systemic hypertension (Final blood pressure in each group had a systolic value above 140mmHg. Diastolic values were below 85 (average around 75. However, standard deviations were on average 9).).
Extra comments	Morphological appearance of aortic valve varied. It was normal in 24 people, bicuspid in 40 people, degenerative in 23 people and rheumatic in 8 people.
Indirectness of population	No indirectness
Interventions	(n=32) Intervention 1: Calcium-channel blockers (CCB) - Nifedipine. Nifedipine 20mg every 12 hours. Duration 7 years. Concurrent medication/care: Not stated.

	Indirectness: No indirectness
	(n=32) Intervention 2: Angiotensin-converting enzyme (ACE) inhibitors - Enalapril . Enalapril 20mg daily. Duration 7 years. Concurrent medication/care: Not stated. Indirectness: No indirectness
	(n=31) Intervention 3: No treatment. No treatment. Duration 7 years. Concurrent medication/care: Not stated. Indirectness: No indirectness
Funding	Academic or government funding (Supported by a grant from the Red de Investigación Cooperativa de las Enfermedades Cardiovasculares from the Instituto de Salud Carlos III, Ministerio de Sanidad y Consumo, Spain )

### RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: NIFEDIPINE versus ENALAPRIL

Protocol outcome 1: All-cause mortality at ≥12 months

- Actual outcome for Primary aortic regurgitation: All-cause mortality at 7 years; Group 1: 1/32, Group 2: 1/32
Risk of bias: All domain - Very high, Selection - High, Blinding - Low, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low,
Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: There are a lot more people with normal morphologic aortic valves in the control group compared to the other groups. There are more women in the control group compared to the others.; Group 1 Number missing: 0, Reason: Nifedipine: 32 people randomised to the treatment. 1 died (sudden death). 7 withdrew due to adverse events. 13 had an aortic valve replacement. Unclear is people reported

were reported as having these events. 1 person is reported to have not received full follow up, their allocated arm was not stated.

; Group 2 Number missing: 0, Reason: Enalapril: 32 people randomised to the treatment. 1 died (Parkinson disease). 3 withdrew due to adverse events. 16 had an aortic valve replacement. Unclear is people reported were reported as having these events. 1 person is reported to have not received full follow up, their allocated arm was not stated.

Protocol outcome 2: Cardiac mortality at ≥12 months

aortic valve replacement. Unclear is people reported

- Actual outcome for Primary aortic regurgitation: Cardiac mortality at 7 years; Group 1: 1/32, Group 2: 0/32
Risk of bias: All domain - Very high, Selection - High, Blinding - Low, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low,
Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: There are a lot more people with normal morphologic aortic valves in the control group compared to the other groups. There are more women in the control group compared to the others.; Group 1 Number missing: 0, Reason: Nifedipine: 32 people randomised to the treatment. 1 died (sudden death). 7 withdrew due to adverse events. 13 had an

were reported as having these events. 1 person is reported to have not received full follow up, their allocated arm was not stated.

; Group 2 Number missing: 0, Reason: Enalapril: 32 people randomised to the treatment. 1 died (Parkinson disease). 3 withdrew due to adverse events. 16 had an aortic valve replacement. Unclear is people reported were reported as having these events. 1 person is reported to have not received full follow up, their allocated arm was not stated.

Protocol outcome 3: Onset of symptoms or progression of NYHA class at ≥12 months

- Actual outcome for Primary aortic regurgitation: Presence of symptoms at 7 years; Group 1: 8/32, Group 2: 10/32
Risk of bias: All domain - Very high, Selection - High, Blinding - Low, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low,
Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: There are a lot more people with normal
morphologic aortic valves in the control group compared to the other groups. There are more women in the control group compared to the others.; Group
1 Number missing: 0, Reason: Nifedipine: 32 people randomised to the treatment. 1 died (sudden death). 7 withdrew due to adverse events. 13 had an
aortic valve replacement. Unclear is people reported

were reported as having these events. 1 person is reported to have not received full follow up, their allocated arm was not stated.

; Group 2 Number missing: 0, Reason: Enalapril: 32 people randomised to the treatment. 1 died (Parkinson disease). 3 withdrew due to adverse events. 16 had an aortic valve replacement. Unclear is people reported were reported as having these events. 1 person is reported to have not received full follow up, their allocated arm was not stated.

Protocol outcome 4: Evidence of HVD progression on imaging (worsening of disease severity) at ≥12 months

- Actual outcome for Primary aortic regurgitation: Left ventricular dysfunction or enlargement on imaging at 7 years; Group 1: 10/32, Group 2: 14/32 Risk of bias: All domain - Very high, Selection - High, Blinding - Low, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: There are a lot more people with normal morphologic aortic valves in the control group compared to the other groups. There are more women in the control group compared to the others.; Group 1 Number missing: 0, Reason: Nifedipine: 32 people randomised to the treatment. 1 died (sudden death). 7 withdrew due to adverse events. 13 had an aortic valve replacement. Unclear is people reported

were reported as having these events. 1 person is reported to have not received full follow up, their allocated arm was not stated.

; Group 2 Number missing: 0, Reason: Enalapril: 32 people randomised to the treatment. 1 died (Parkinson disease). 3 withdrew due to adverse events. 16 had an aortic valve replacement. Unclear is people reported were reported as having these events. 1 person is reported to have not received full follow up, their allocated arm was not stated.

Protocol outcome 5: Need for heart valve intervention at ≥12 months

- Actual outcome for Primary aortic regurgitation: Aortic valve replacement at 7 years; Group 1: 13/32, Group 2: 16/32
Risk of bias: All domain - Very high, Selection - High, Blinding - Low, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low,
Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: There are a lot more people with normal morphologic aortic valves in the control group compared to the other groups. There are more women in the control group compared to the others.; Group 1 Number missing: 0, Reason: Nifedipine: 32 people randomised to the treatment. 1 died (sudden death). 7 withdrew due to adverse events. 13 had an aortic valve replacement. Unclear is people reported

were reported as having these events. 1 person is reported to have not received full follow up, their allocated arm was not stated.

; Group 2 Number missing: 0, Reason: Enalapril: 32 people randomised to the treatment. 1 died (Parkinson disease). 3 withdrew due to adverse events. 16 had an aortic valve replacement. Unclear is people reported were reported as having these events. 1 person is reported to have not received full follow up, their allocated arm was not stated.

Protocol outcome 6: Withdrawal due to adverse events at ≥12 months

- Actual outcome for Primary aortic regurgitation: Withdrawal due to adverse events at 7 years; Group 1: 7/32, Group 2: 3/32; Comments: Nifedipine: 7 developed either headache, flushing, oedema, epigastric pain or a combination of these. Enalapril: 2 developed cough, 1 developed hypotension Risk of bias: All domain - Very high, Selection - High, Blinding - Low, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: There are a lot more people with normal morphologic aortic valves in the control group compared to the other groups. There are more women in the control group compared to the others.; Group 1 Number missing: 0, Reason: Nifedipine: 32 people randomised to the treatment. 1 died (sudden death). 7 withdrew due to adverse events. 13 had an aortic valve replacement. Unclear is people reported

were reported as having these events. 1 person is reported to have not received full follow up, their allocated arm was not stated.

; Group 2 Number missing: 0, Reason: Enalapril: 32 people randomised to the treatment. 1 died (Parkinson disease). 3 withdrew due to adverse events. 16 had an aortic valve replacement. Unclear is people reported were reported as having these events. 1 person is reported to have not received full follow up, their allocated arm was not stated.

RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: NIFEDIPINE versus NO TREATMENT

Protocol outcome 1: All-cause mortality at ≥12 months

- Actual outcome for Primary aortic regurgitation: All-cause mortality at 7 years; Group 1: 1/32, Group 2: 1/31
Risk of bias: All domain - Very high, Selection - High, Blinding - Low, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: There are a lot more people with normal morphologic aortic valves in the control group compared to the others.: Group

1 Number missing: 0, Reason: Nifedipine: 32 people randomised to the treatment. 1 died (sudden death). 7 withdrew due to adverse events. 13 had an aortic valve replacement. Unclear is people reported

were reported as having these events. 1 person is reported to have not received full follow up, their allocated arm was not stated.

; Group 2 Number missing: 0, Reason: Placebo: 31 people randomised to the treatment. 1 died (of heart failure). 12 had an aortic valve replacement. Unclear is people reported were reported as having

these events. 1 person is reported to have not received full follow up, their allocated arm was not stated.

Protocol outcome 2: Cardiac mortality at ≥12 months

- Actual outcome for Primary aortic regurgitation : Cardiac mortality at 7 years; Group 1: 1/32, Group 2: 1/31

Risk of bias: All domain - Very high, Selection - High, Blinding - Low, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: There are a lot more people with normal morphologic aortic valves in the control group compared to the other groups. There are more women in the control group compared to the others.; Group 1 Number missing: 0, Reason: Nifedipine: 32 people randomised to the treatment. 1 died (sudden death). 7 withdrew due to adverse events. 13 had an aortic valve replacement. Unclear is people reported

were reported as having these events. 1 person is reported to have not received full follow up, their allocated arm was not stated.

; Group 2 Number missing: 0, Reason: Placebo: 31 people randomised to the treatment. 1 died (of heart failure). 12 had an aortic valve replacement. Unclear is people reported were reported as having

these events. 1 person is reported to have not received full follow up, their allocated arm was not stated.

Protocol outcome 3: Onset of symptoms or progression of NYHA class at ≥12 months

- Actual outcome for Primary aortic regurgitation : Presence of symptoms at 7 years; Group 1: 8/32, Group 2: 8/31

Risk of bias: All domain - Very high, Selection - High, Blinding - Low, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: There are a lot more people with normal morphologic aortic valves in the control group compared to the other groups. There are more women in the control group compared to the others.; Group 1 Number missing: 0, Reason: Nifedipine: 32 people randomised to the treatment. 1 died (sudden death). 7 withdrew due to adverse events. 13 had an aortic valve replacement. Unclear is people reported

were reported as having these events. 1 person is reported to have not received full follow up, their allocated arm was not stated.

; Group 2 Number missing: 0, Reason: Placebo: 31 people randomised to the treatment. 1 died (of heart failure). 12 had an aortic valve replacement. Unclear is people reported were reported as having

these events. 1 person is reported to have not received full follow up, their allocated arm was not stated.

Protocol outcome 4: Evidence of HVD progression on imaging (worsening of disease severity) at ≥12 months

- Actual outcome for Primary aortic regurgitation: Left ventricular dysfunction or enlargement on imaging at 7 years; Group 1: 10/32, Group 2: 10/31 Risk of bias: All domain - Very high, Selection - High, Blinding - Low, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: There are a lot more people with normal morphologic aortic valves in the control group compared to the other groups. There are more women in the control group compared to the others.; Group 1 Number missing: 0, Reason: Nifedipine: 32 people randomised to the treatment. 1 died (sudden death). 7 withdrew due to adverse events. 13 had an aortic valve replacement. Unclear is people reported

were reported as having these events. 1 person is reported to have not received full follow up, their allocated arm was not stated.

; Group 2 Number missing: 0, Reason: Placebo: 31 people randomised to the treatment. 1 died (of heart failure). 12 had an aortic valve replacement. Unclear is people reported were reported as having

these events. 1 person is reported to have not received full follow up, their allocated arm was not stated.

Protocol outcome 5: Need for heart valve intervention at ≥12 months

- Actual outcome for Primary aortic regurgitation: Aortic valve replacement at 7 years; Group 1: 13/32, Group 2: 12/31
Risk of bias: All domain - Very high, Selection - High, Blinding - Low, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: There are a lot more people with normal morphologic aortic valves in the control group compared to the other groups. There are more women in the control group compared to the others.; Group 1 Number missing: 0, Reason: Nifedipine: 32 people randomised to the treatment. 1 died (sudden death). 7 withdrew due to adverse events. 13 had an aortic valve replacement. Unclear is people reported

were reported as having these events. 1 person is reported to have not received full follow up, their allocated arm was not stated.

; Group 2 Number missing: 0, Reason: Placebo: 31 people randomised to the treatment. 1 died (of heart failure). 12 had an aortic valve replacement. Unclear is people reported were reported as having

these events. 1 person is reported to have not received full follow up, their allocated arm was not stated.

Protocol outcome 6: Withdrawal due to adverse events at ≥12 months

- Actual outcome for Primary aortic regurgitation: Withdrawal due to adverse events at 7 years; Group 1: 7/32, Group 2: 0/31; Comments: Nifedipine: 7 developed either headache, flushing, oedema, epigastric pain or a combination of these.

Risk of bias: All domain - Very high, Selection - High, Blinding - Low, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: There are a lot more people with normal morphologic aortic valves in the control group compared to the other groups. There are more women in the control group compared to the others.; Group 1 Number missing: 0, Reason: Nifedipine: 32 people randomised to the treatment. 1 died (sudden death). 7 withdrew due to adverse events. 13 had an aortic valve replacement. Unclear is people reported

were reported as having these events. 1 person is reported to have not received full follow up, their allocated arm was not stated.

; Group 2 Number missing: 0, Reason: Placebo: 31 people randomised to the treatment. 1 died (of heart failure). 12 had an aortic valve replacement. Unclear is people reported were reported as having these events. 1 person is reported to have not received full follow up, their allocated arm was not stated.

# RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: ENALAPRIL versus NO TREATMENT

Protocol outcome 1: All-cause mortality at ≥12 months

- Actual outcome for Primary aortic regurgitation: All-cause mortality at 7 years; Group 1: 1/32, Group 2: 1/31
Risk of bias: All domain - Very high, Selection - High, Blinding - Low, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low,
Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: There are a lot more people with normal
morphologic aortic valves in the control group compared to the other groups. There are more women in the control group compared to the others.; Group
1 Number missing: 0, Reason: Enalapril: 32 people randomised to the treatment. 1 died (Parkinson disease). 3 withdrew due to adverse events. 16 had an
aortic valve replacement. Unclear is people reported were reported as having these events. 1 person is reported to have not received full follow up, their
allocated arm was not stated.

; Group 2 Number missing: 0, Reason: Placebo: 31 people randomised to the treatment. 1 died (of heart failure). 12 had an aortic valve replacement. Unclear is people reported were reported as having these events. 1 person is reported to have not received full follow up, their allocated arm was not stated.

Protocol outcome 2: Cardiac mortality at ≥12 months

- Actual outcome for Primary aortic regurgitation : Cardiac mortality at 7 years; Group 1: 0/32, Group 2: 1/31

Risk of bias: All domain - Very high, Selection - High, Blinding - Low, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: There are a lot more people with normal morphologic aortic valves in the control group compared to the other groups. There are more women in the control group compared to the others.; Group 1 Number missing: 0, Reason: Enalapril: 32 people randomised to the treatment. 1 died (Parkinson disease). 3 withdrew due to adverse events. 16 had an aortic valve replacement. Unclear is people reported were reported as having these events. 1 person is reported to have not received full follow up, their allocated arm was not stated.

; Group 2 Number missing: 0, Reason: Placebo: 31 people randomised to the treatment. 1 died (of heart failure). 12 had an aortic valve replacement. Unclear is people reported were reported as having

these events. 1 person is reported to have not received full follow up, their allocated arm was not stated.

Protocol outcome 3: Onset of symptoms or progression of NYHA class at ≥12 months

- Actual outcome for Primary aortic regurgitation : Presence of symptoms at 7 years; Group 1: 10/32, Group 2: 8/31

Risk of bias: All domain - Very high, Selection - High, Blinding - Low, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: There are a lot more people with normal morphologic aortic valves in the control group compared to the other groups. There are more women in the control group compared to the others.; Group 1 Number missing: 0, Reason: Enalapril: 32 people randomised to the treatment. 1 died (Parkinson disease). 3 withdrew due to adverse events. 16 had an aortic valve replacement. Unclear is people reported were reported as having these events. 1 person is reported to have not received full follow up, their allocated arm was not stated.

; Group 2 Number missing: 0, Reason: Placebo: 31 people randomised to the treatment. 1 died (of heart failure). 12 had an aortic valve replacement. Unclear is people reported were reported as having

these events. 1 person is reported to have not received full follow up, their allocated arm was not stated.

Protocol outcome 4: Evidence of HVD progression on imaging (worsening of disease severity) at ≥12 months

- Actual outcome for Primary aortic regurgitation: Left ventricular dysfunction or enlargement on imaging at 7 years; Group 1: 14/32, Group 2: 10/31 Risk of bias: All domain - Very high, Selection - High, Blinding - Low, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: There are a lot more people with normal

morphologic aortic valves in the control group compared to the other groups. There are more women in the control group compared to the others.; Group 1 Number missing: 0, Reason: Enalapril: 32 people randomised to the treatment. 1 died (Parkinson disease). 3 withdrew due to adverse events. 16 had an aortic valve replacement. Unclear is people reported were reported as having these events. 1 person is reported to have not received full follow up, their allocated arm was not stated.

; Group 2 Number missing: 0, Reason: Placebo: 31 people randomised to the treatment. 1 died (of heart failure). 12 had an aortic valve replacement. Unclear is people reported were reported as having these events. 1 person is reported to have not received full follow up, their allocated arm was not stated.

Protocol outcome 5: Need for heart valve intervention at ≥12 months

- Actual outcome for Primary aortic regurgitation: Aortic valve replacement at 7 years; Group 1: 16/32, Group 2: 12/31
  Risk of bias: All domain Very high, Selection High, Blinding Low, Incomplete outcome data High, Outcome reporting Low, Measurement Low,
  Crossover Low, Subgroups Low, Other 1 Low; Indirectness of outcome: No indirectness; Baseline details: There are a lot more people with normal
  morphologic aortic valves in the control group compared to the other groups. There are more women in the control group compared to the others.; Group
  1 Number missing: 0, Reason: Enalapril: 32 people randomised to the treatment. 1 died (Parkinson disease). 3 withdrew due to adverse events. 16 had an
  aortic valve replacement. Unclear is people reported were reported as having these events. 1 person is reported to have not received full follow up, their
  allocated arm was not stated.
- ; Group 2 Number missing: 0, Reason: Placebo: 31 people randomised to the treatment. 1 died (of heart failure). 12 had an aortic valve replacement. Unclear is people reported were reported as having these events. 1 person is reported to have not received full follow up, their allocated arm was not stated.

Protocol outcome 6: Withdrawal due to adverse events at ≥12 months

- Actual outcome for Primary aortic regurgitation : Withdrawal due to adverse events at 7 years; Group 1: 3/32, Group 2: 0/31; Comments: Enalapril: 2 developed cough, 1 developed hypotension

Risk of bias: All domain - Very high, Selection - High, Blinding - Low, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: There are a lot more people with normal morphologic aortic valves in the control group compared to the other groups. There are more women in the control group compared to the others.; Group 1 Number missing: 0, Reason: Enalapril: 32 people randomised to the treatment. 1 died (Parkinson disease). 3 withdrew due to adverse events. 16 had an aortic valve replacement. Unclear is people reported were reported as having these events. 1 person is reported to have not received full follow up, their allocated arm was not stated.

; Group 2 Number missing: 0, Reason: Placebo: 31 people randomise Unclear is people reported were reported as having these events. 1 person is reported to have not received full follow up,	ed to the treatment. 1 died (of heart failure). 12 had an aortic valve replacement. their allocated arm was not stated.
Protocol outcomes not reported by the study	Quality of life at 6 months; Quality of life at ≥12 months; Exercise tolerance at ≥12 months; Withdrawal due to adverse events at 6 months

Study	Hansson 2017 <sup>66</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=40)
Countries and setting	Conducted in Denmark; Setting: Outpatient follow up
Line of therapy	1st line
Duration of study	Intervention + follow up: 22 weeks (5 months)
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: Echocardiography and cardiac MR
Stratum	Primary aortic [including bicuspid] stenosis
Subgroup analysis within study	Not applicable
Inclusion criteria	Asymptomatic aortic stenosis with an aortic valve area ≤1.2cm² or transaortic maximal velocity ≥3.0m/s and sinus rhythm with an HR ≥60/min.
Exclusion criteria	Ongoing treatment with beta blockers, significant aortic valve regurgitation (vena contracta ≥5mm), or ischaemic heart disease evaluated by symptoms or signs of myocardial ischaemia (i.e. angina pectoris, abnormal echocardiography, wall motion abnormalities). In addition, people with previous coronary angiography proving a ≥70% luminal stenosis were excluded.
Recruitment/selection of patients	Recruitment from outpatient clinics at 3 centres between August 2013 and April 2016.
Age, gender and ethnicity	Age - Mean (SD): 70.0 (5.1). Gender (M:F): 24:14. Ethnicity: Not stated
Further population details	1. Age: Mixed (Based on mean age and standard deviation). 2. Disease mechanism for aortic and mitral stenosis: Not stated / Unclear (States that 7 people had bicuspid aortic valves, but no other information.). 3. Presence vs. absence of uncontrolled systemic hypertension (140/85 mmHg) at the end of trial intervention despite medication: Mixed (Mean blood pressure at follow up metoprolol: 136/79 (13/8). Placebo: 140/81 (12/7).
Indirectness of population	No indirectness
Interventions	(n=20) Intervention 1: Beta blockers - Metoprolol. Extended-release metoprolol from 50mg up to a target dose of 200mg or maximal dose without symptoms. Achieved during a 6-week uptitration period. Duration 5 months. Concurrent medication/care: Not stated. Indirectness: No indirectness
	(n=20) Intervention 2: Placebo. Placebo. Duration 5 months. Concurrent

Ni Eli	cademic or government funding (Funded by the Lundbeck foundation, the Arvid lilssons Foundation, the Health Research Fund of Central Denmark Region, Karen lise Jensens Foundation, and Snedkermester Sophus Jacobsen and Hustru Astrid acobsens Foundation.)

### RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: METOPROLOL versus PLACEBO

Protocol outcome 1: Quality of life at 6 months

- Actual outcome for Primary aortic [including bicuspid] stenosis: Minnesota living with heart failure questionnaire at 5 months; Group 1: mean 5 (SD 14); n=19, Group 2: mean -1 (SD 4); n=19; Minnesota living with heart failure questionnaire 0-105 Top=High is poor outcome; Comments: Baseline (median (IQR)) metoprolol: 3 (1-6). Baseline placebo: 4 (2-8). Final metoprolol: 5 (2-9). Final placebo: 3 (0-8).

Risk of bias: All domain - Very high, Selection - Very high, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: Generally people in the placebo group had more comorbidities (hypertension and diabetes mellitus) and used more medication than the metoprolol group. However, this is based on a very small number of people.; Group 1 Number missing: 1, Reason: 1 excluded due to having an LVEF <50% after randomisation.; Group 2 Number missing: 1, Reason: 1 excluded due to having an LVEF <50% after randomisation.

### Protocol outcome 2: Exercise tolerance at ≥12 months

- Actual outcome for Primary aortic [including bicuspid] stenosis : 6 minute walk test distance, m at 5 months; Group 1: mean 2 m (SD 46); n=19, Group 2: mean 14 m (SD 49); n=19; Comments: Baseline metoprolol: 543 (46)m. Baseline placebo: 538 (36)m. Final metoprolol: 546 (48)m. Final placebo: 550 (49)m.

Risk of bias: All domain - Very high, Selection - Very high, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: Generally people in the placebo group had more comorbidities (hypertension and diabetes mellitus) and used more medication than the metoprolol group. However, this is based on a very small number of people.; Group 1 Number missing: 1, Reason: 1 excluded due to having an LVEF <50% after randomisation.; Group 2 Number missing: 1, Reason: 1 excluded due to having an LVEF <50% after randomisation.

Protocol outcome 3: Withdrawal due to adverse events at 6 months

- Actual outcome for Primary aortic [including bicuspid] stenosis : Withdrawal or dose reduction due to adverse events at 5 months; Group 1: 4/19, Group 2: 2/19

Risk of bias: All domain - Very high, Selection - Very high, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: Serious indirectness, Comments: Includes people who had a dose reduction and then continued the trial with no additional adverse events; Baseline details: Generally people in the placebo group had more comorbidities (hypertension and diabetes mellitus) and used more medication than the metoprolol group. However, this is based on a very small number of people.; Group 1 Number missing: 1, Reason: 1 excluded due to having an LVEF <50% after randomisation.; Group 2 Number missing: 1, Reason: 1 excluded due

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to having an LVEF <50% after randomisation.	
Protocol outcomes not reported by the study	All-cause mortality at ≥12 months; Cardiac mortality at ≥12 months; Quality of life at ≥12 months; Onset of symptoms or progression of NYHA class at ≥12 months; Evidence of HVD progression on imaging (worsening of disease severity) at ≥12 months; Need for heart valve intervention at ≥12 months; Withdrawal due to adverse events at ≥12 months

Study
Study type
Number o
Countries
Line of the
Duration of
Method of
Stratum
Subgroup
Inclusion
Exclusion
Docruitmo

Study	Marcotte 1997 <sup>104</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=23)
Countries and setting	Conducted in Canada; Setting: Outpatient follow up
Line of therapy	1st line
Duration of study	Intervention + follow up: 1 year
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: Echocardiography (Hewlett-Packard Sonos 1500 ultrasonograph) with 2.5 and 3.5 mHz transducers
Stratum	Primary mitral regurgitation
Subgroup analysis within study	Not applicable
Inclusion criteria	Asymptomatic adults (18 to 75 years) with at least moderate, organic (degenerative, rheumatic, postinfectious or congenital), isolated mitral regurgitation (grade 3+ or more). People were in sinus rhythm, taking no cardiovascular medication at the time of enrolment. They needed to have had a good quality echocardiogram showing a maximal MR colour jet area >4cm² or greater than 25% of the LA area in at least two different views, and normal LVEF (>60%).
Exclusion criteria	Clinically documented coronary artery disease (including angina, prior myocardial infarction and prior revascularisation); mitral stenosis (valve area less than 2.5cm²); significant ventricular or atrial arrhythmia (including atrial fibrillation); significant aortic valve disease (either moderate to severe aortic regurgitation or the presence of aortic stenosis, defined as a valve area less than 2.0cm²); hypertension under therapy before randomisation or untreated hypertension with a DBP >90mmHg or hypotension with an SBP <90mmHg; chronic renal failure; contraindication to receiving ACE inhibitors; abnormal LV systolic function by echocardiography (<60%); severe LV dilation; or resting or stress-induced regional wall motion abnormalities.
Recruitment/selection of patients	10,054 people underwent echocardiography at the author's institution. 248 satisfied the echocardiographic inclusion criteria. 211 refused to participate in the study, were judged to be symptomatic, had undergone mitral valve surgery, or presented clinical or echocardiographic exclusion criteria. 37 attended a screening visit, of which 5 refused to participate and 9 were found to have clinical or echocardiographic exclusion factors.
Age, gender and ethnicity	Age - Other: 53.3 (standard error: 2.4). Gender (M:F): 16:7. Ethnicity: Not stated

Further population details	1. Age: <75 years (Mean age: 53.3 (standard error: 2.4)). 2. Disease mechanism for aortic and mitral stenosis: Not applicable 3. Presence vs. absence of uncontrolled systemic hypertension (140/85 mmHg) at the end of trial intervention despite medication: Not stated / Unclear (None at the start of the study, but not reported at the end).
Indirectness of population	Serious indirectness: Population may include people with congenital mitral regurgitation
Interventions	<ul> <li>(n=12) Intervention 1: Angiotensin-converting enzyme (ACE) inhibitors - Lisinopril. Lisinopril 5mg for two weeks, then doubled every two weeks until the maximal dose of 20mg a day was reached or they developed symptoms of hypotension, in which the dose was titrated to the maximal tolerable dose. Duration 1 year. Concurrent medication/care: No other cardiovascular medications. Indirectness: No indirectness</li> <li>(n=11) Intervention 2: Placebo. Placebo. Duration 1 year. Concurrent medication/care: No other cardiovascular medications. Indirectness: No indirectness</li> </ul>
Funding	Study funded by industry (Supported by a grant from Merck Frosst Canada inc.)

# RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: LISINOPRIL versus PLACEBO

## Protocol outcome 1: All-cause mortality at ≥12 months

- Actual outcome for Primary mitral regurgitation: Death at 1 year; Group 1: 0/6, Group 2: 0/10

Risk of bias: All domain - Very high, Selection - High, Blinding - Low, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: Exercise time different between the two groups (lisinopril = 581 (37), placebo = 637 (56)). This may affect the results of exercise time.; Group 1 Number missing: 6, Reason: 4 withdrew due to adverse events. 2 additional people were lost due to need for surgical intervention or urgent medical therapy.; Group 2 Number missing: 1, Reason: 1 withdrawal due to adverse events.

## Protocol outcome 2: Cardiac mortality at ≥12 months

- Actual outcome for Primary mitral regurgitation: Death at 1 year; Group 1: 0/6, Group 2: 0/10

Risk of bias: All domain - Very high, Selection - High, Blinding - Low, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: Exercise time different between the two groups (lisinopril = 581 (37), placebo = 637 (56)). This may affect the results of exercise time.; Group 1 Number missing: 6, Reason: 4 withdrew due to adverse events. 2 additional people were lost due to need for surgical intervention or urgent medical therapy.; Group 2 Number missing: 1, Reason: 1 withdrawal due to adverse events.

### Protocol outcome 3: Quality of life at 6 months

- Actual outcome for Primary mitral regurgitation: Life quality index at 6 months; Group 1: mean 0.2 (SD 0.73); n=6, Group 2: mean 0.4 (SD 0.95); n=10; Life quality index 1-6 Top=High is good outcome; Comments: Life quality index is a measurement for asymptomatic hypertensive people where they rate their energy level on the scale from 1 (no energy) to 6 (full of energy). Reports standard error. Lisinopril 6 months: +0.2 (0.3). Placebo 6 months: +0.3 (0.3). Lisinopril baseline: 4.6 (0.4).

Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - High, Outcome reporting - Low, Measurement - High, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: Exercise time different between the two groups (lisinopril = 581 (37), placebo = 637 (56)). This may affect the results of exercise time.; Group 1 Number missing: 6, Reason: 4 withdrew due to adverse events. 2 additional people were lost due to need for surgical intervention or urgent medical therapy.; Group 2 Number missing: 1, Reason: 1 withdrawal due to adverse events.

### Protocol outcome 4: Quality of life at ≥12 months

- Actual outcome for Primary mitral regurgitation: Life quality index at 1 year; Group 1: mean 0.3 (SD 0.73); n=6, Group 2: mean 0.4 (SD 0.95); n=10; Life quality index 1-6 Top=High is good outcome; Comments: Life quality index is a measurement for asymptomatic hypertensive people where they rate their energy level on the scale from 1 (no energy) to 6 (full of energy). Reports standard error. Lisinopril 1 year: +0.3 (0.3). Placebo 1 year: +0.4 (0.3). Lisinopril baseline: 4.6 (0.4). Placebo baseline: 4.8 (0.4).

Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - High, Outcome reporting - Low, Measurement - High, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: Exercise time different between the two groups (lisinopril = 581 (37), placebo = 637 (56)). This may affect the results of exercise time.; Group 1 Number missing: 6, Reason: 4 withdrew due to adverse events. 2 additional people were lost due to need for surgical intervention or urgent medical therapy.; Group 2 Number missing: 1, Reason: 1 withdrawal due to adverse events.

#### Protocol outcome 5: Exercise tolerance at ≥12 months

- Actual outcome for Primary mitral regurgitation: Treadmill exercise time (Bruce protocol) at 1 year; Group 1: mean 39 seconds (SD 61.2); n=6, Group 2: mean 18 seconds (SD 66.4); n=10; Comments: Reported with standard errors. Lisinopril 1 year: 39 (25)s. Placebo 1 year: 18 (21)s. Lisinopril baseline: 581 (37)s. Placebo baseline: 637 (56)s.

Risk of bias: All domain - Very high, Selection - Very high, Blinding - Low, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: Exercise time different between the two groups (lisinopril = 581 (37), placebo = 637 (56)). This may affect the results of exercise time.; Group 1 Number missing: 6, Reason: 4 withdrew due to adverse events. 2 additional people were lost due to need for surgical intervention or urgent medical therapy.; Group 2 Number missing: 1, Reason: 1 withdrawal due to adverse events.

### Protocol outcome 6: Withdrawal due to adverse events at ≥12 months

- Actual outcome for Primary mitral regurgitation: Withdrawal due to adverse events at 12 months; Group 1: 4/10, Group 2: 1/11; Comments: The study reports withdrawal due to adverse events as a whole rather than when people withdrew from the study. Therefore, the results are being reported for the longest time period possible only.

Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low,

	utcome: No indirectness; Baseline details: Exercise time different between the two e results of exercise time.; Group 1 Number missing: 2, Reason: 2 people were lost oup 2 Number missing: 0
Protocol outcomes not reported by the study	Onset of symptoms or progression of NYHA class at ≥12 months; Evidence of HVD progression on imaging (worsening of disease severity) at ≥12 months; Need for heart valve intervention at ≥12 months; Withdrawal due to adverse events at 6 months

Study	Roberts 2018 <sup>129</sup>
Study type	RCT (Patient randomised; Crossover: 0 days)
Number of studies (number of participants)	1 (n=46)
Countries and setting	Conducted in New Zealand; Setting: Secondary care
Line of therapy	1st line
Duration of study	Intervention + follow up: Maximum 7 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: Echocardiography (people were recruited from a clinical echocardiography database).
Stratum	Primary aortic regurgitation
Subgroup analysis within study	Not applicable
Inclusion criteria	People with moderate to severe chronic aortic regurgitation (two or more of the following present on echocardiography: aortic regurgitant central jet width >25% of left ventricular outflow tract, vena contracta width >0.3cm, presence of early diastolic flow reversal in the descending aorta, pressure half time or aortic regurgitant velocity <500ms, and left ventricular end diastolic dimension or volume above the normal reference range) and normal left ventricular systolic function (ejection fraction >50%).
Exclusion criteria	Age <18 or >80 years; contraindications to cardiac magnetic resonance imaging; inability to complete study exercise protocol or procedures; other documented significant cardiac diseases; contraindications to study medications or withdrawal of usual antihypertensive medications.
Recruitment/selection of patients	Recruited from a clinical echocardiography database at Auckland City Hospital.
Age, gender and ethnicity	Age - Mean (SD): 51.0 (14.1). Gender (M:F): 14:3. Ethnicity: Not stated
Further population details	1. Age: <75 years 2. Disease mechanism for aortic and mitral stenosis: Not applicable 3. Presence vs. absence of uncontrolled systemic hypertension (140/85 mmHg) at the end of trial intervention despite medication: Absence of uncontrolled systemic hypertension (Blood pressure ranged between 117-118/63-69.).
Indirectness of population	No indirectness
Interventions	(n=17) Intervention 1: Angiotensin-II receptor antagonists (ARBs) - Losartan . Losartan up-titrated to a maximum of 100mg per day. Duration 1-3 weeks. Concurrent medication/care: People were allowed to use normal antihypertensive medicines, which were then downtitrated or withdrawn completely while taking the drug. Indirectness: Serious indirectness; Indirectness comment: Inadequate

	duration of treatment  (n=17) Intervention 2: Beta blockers - Metoprolol . Metoprolol CR to a maximum dose of 190mg. Duration 1-3 weeks. Concurrent medication/care: People were allowed to use normal antihypertensive medicines, which were then downtitrated or withdrawn completely while taking the drug. Indirectness: Serious indirectness; Indirectness comment: Inadequate duration of treatment
Funding	Academic or government funding (Funded by the Health Research Council of New Zealand)
RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COM  Protocol outcome 1: Exercise tolerance at ≥12 months	IPARISON: LOSARTAN versus METOPROLOL

- Actual outcome for Primary aortic regurgitation: Exercise work rate (Watts) - using an ergometer at 3 weeks; Group 1: mean 29 Watts (SD 6); n=17, Group 2: mean 29 Watts (SD 8); n=17; Comments: Not provided with baseline values

Risk of bias: All domain - Very high, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - High, Subgroups - Low, Other 1 - Low; Indirectness of outcome: Serious indirectness, Comments: Time period for treatment was less than 1 month; Group 1 Number missing: 0; Group 2 Number missing: 0

Protocol outcomes not reported by the study

All-cause mortality at ≥12 months; Cardiac mortality at ≥12 months; Quality of life at ≥12 months; Quality of life at 6 months; Onset of symptoms or progression of NYHA class at ≥12 months; Evidence of HVD progression on imaging (worsening of disease severity) at ≥12 months; Need for heart valve intervention at ≥12 months; Withdrawal due to adverse events at 6 months; Withdrawal due to adverse events at ≥12 months

4 MIDs used to assess imprecision were ±4.0

leart valve disease: DRAFT FOR CONSULTATION

Study (subsidiary papers)	SALTIRE trial: Cowell 2005 <sup>36</sup> (Houslay 2006 <sup>74</sup> )
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=155)
Countries and setting	Conducted in United Kingdom; Setting: Outpatient follow up
Line of therapy	1st line
Duration of study	Intervention + follow up: 25 months
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: Echocardiography
Stratum	Primary aortic [including bicuspid] stenosis
Subgroup analysis within study	Not applicable
Inclusion criteria	People older than 18 years with calcific aortic stenosis, an aortic-jet velocity of at least 2.5m/s, and aortic-valve calcification on echocardiography.
Exclusion criteria	Child-bearing potential without contraception, active or chronic liver disease, a history of alcohol or drug abuse, severe mitral-valve stenosis (mitral-valve area <1cm²), severe mitral or aortic regurgitation, left ventricular dysfunction (EF <35%), a planned aortic-valve replacement, intolerance of statins, statin therapy or a potential benefit from statin therapy (according to the treating physician), a baseline serum total cholesterol concentration of less than 150mg/dL (4.0mmol/L), and presence of a permanent pacemaker or cardiodefibrillator.
Recruitment/selection of patients	Recruited from eight centers.
Age, gender and ethnicity	Age - Mean (SD): 68 (10.5). Gender (M:F): 140:15. Ethnicity: Not stated
Further population details	1. Age: Mixed (Confidence intervals fall either side of 75 years.). 2. Disease mechanism for aortic and mitral stenosis: Calcific 3. Presence vs. absence of uncontrolled systemic hypertension (140/85 mmHg) at the end of trial intervention despite medication: Not stated / Unclear (Baseline atorvastatin: 144/82 (18/10). Baseline placebo: 144.81 (21/12). No final value stated.).
Indirectness of population	No indirectness
Interventions	(n=77) Intervention 1: Statins - Atorvastatin . Atorvastatin 80mg once a day. Duration 25 months. Concurrent medication/care: 43 taking aspirin, 12 taking ACE inhibitors, 21 taking beta-blockers, 8 taking warfarin. Otherwise not stated. Indirectness: No indirectness
	(n=78) Intervention 2: Placebo. Placebo. Duration 25 months. Concurrent medication/care: 40 took aspirin, 14 took ACE inhibitors, 27 took beta-blockers, 12

	took warfarin. Otherwise not stated. Indirectness: No indirectness
Funding	Funding not stated

### RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: ATORVASTATIN versus PLACEBO

Protocol outcome 1: Cardiac mortality at ≥12 months

- Actual outcome for Primary aortic [including bicuspid] stenosis: Death from cardiovascular causes at 25 months; Group 1: 3/77, Group 2: 3/78
Risk of bias: All domain - Low, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcome 2: Onset of symptoms or progression of NYHA class at ≥12 months

- Actual outcome for Primary aortic [including bicuspid] stenosis: Hospitalisation for severe aortic stenosis at 25 months; Group 1: 3/77, Group 2: 5/78
Risk of bias: All domain - Low, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Group 1 Number missing:

### Protocol outcome 3: Need for heart valve intervention at ≥12 months

- Actual outcome for Primary aortic [including bicuspid] stenosis: Aortic valve replacement at 25 months; Group 1: 11/77, Group 2: 19/78
Risk of bias: All domain - Low, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: ; Group 2 Number missing:

#### Protocol outcome 4: Withdrawal due to adverse events at ≥12 months

- Actual outcome for Primary aortic [including bicuspid] stenosis: Withdrawal due to adverse events at 25 months; Group 1: 7/77, Group 2: 4/78
Risk of bias: All domain - Low, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the study	All-cause mortality at ≥12 months; Quality of life at 6 months; Quality of life at ≥12
	months; Evidence of HVD progression on imaging (worsening of disease severity)
	at ≥12 months; Exercise tolerance at ≥12 months; Withdrawal due to adverse
	events at 6 months

Study	Sampaio 2005 <sup>143</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=47)
Countries and setting	Conducted in Brazil; Setting: Outpatient follow up
Line of therapy	1st line
Duration of study	Intervention + follow up: 1 year
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: Echocardiography
Stratum	Primary mitral regurgitation
Subgroup analysis within study	Not applicable
Inclusion criteria	People with moderate to severe mitral regurgitation according to echocardiographic criteria: mitral regurgitation jet area >40% of the left atrium or no evidence of leaflet coaptation; absolute MR jet area >8cm² associated with left chamber dilation; MR jet into pulmonary veins.
Exclusion criteria	Atrial fibrillation; systolic blood pressure <100 or >160 mmHg; receiving therapy with other vasodilators.
Recruitment/selection of patients	No additional information given
Age, gender and ethnicity	Age - Mean (SD): 39 (15) years. Gender (M:F): 27:20. Ethnicity: Not stated
Further population details	1. Age: <75 years (Mean age: 39 (15) years). 2. Disease mechanism for aortic and mitral stenosis: Not applicable 3. Presence vs. absence of uncontrolled systemic hypertension (140/85 mmHg) at the end of trial intervention despite medication: Absence of uncontrolled systemic hypertension (At 12 months the mean systolic blood pressure for the enalapril arm: 122 (13)mmHg. Placebo arm: 126 (12)mmHg. Mean diastolic blood pressure for the enalapril arm: 78 (9). Placebo arm: 79 (8).).
Extra comments	. At the start of the study, 20 people were NYHA class I, 17 people were NYHA class II.
Indirectness of population	Serious indirectness
Interventions	(n=27) Intervention 1: Angiotensin-converting enzyme (ACE) inhibitors - Enalapril . Enalapril 5mg twice a day, titrated up to the maximal tolerated dose of at most 20mg twice a day with increases to 10mg and then 20mg at 2 weekly intervals if systolic blood pressure remained >100mmHg. Duration 1 year. Concurrent medication/care: Not receiving therapy with any other vasodilators.  (n=27) Intervention 2: Placebo. Placebo. Duration 1 year. Concurrent

	medication/care: Not receiving therapy with any other vasodilators. Indirectness: No indirectness
Funding	Academic or government funding (Received grants from the E.J. Zerbini foundation, São Paulo, Brazil)

## RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: ENALAPRIL versus PLACEBO

Protocol outcome 1: Onset of symptoms or progression of NYHA class at ≥12 months

- Actual outcome for Primary mitral regurgitation: Onset of symptoms or progression of NYHA class at 12 months; Group 1: 0/26, Group 2: 4/22 Risk of bias: All domain - Very high, Selection - Very high, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: People in the placebo arm appear to have more severe valve disease and worse exercise tolerance than the enalapril arm.; Group 1 Number missing: 5, Reason: 5 withdrew due to nonadherence; Group 2 Number missing: 1, Reason: 1 withdrew due to nonadherence

### Protocol outcome 2: Need for heart valve intervention at ≥12 months

- Actual outcome for Primary mitral regurgitation: Need for heart valve intervention at 12 months; Group 1: 0/26, Group 2: 1/22 Risk of bias: All domain - Very high, Selection - Very high, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: People in the placebo arm appear to have more severe valve disease and worse exercise tolerance than the enalapril arm.; Group 1 Number missing: 5, Reason: 5 withdrew due to nonadherence; Group 2 Number missing: 1, Reason: 1 withdrew due to nonadherence

#### Protocol outcome 3: Exercise tolerance at ≥12 months

- Actual outcome for Primary mitral regurgitation: Oxygen uptake at peak exercise (mL/min) at 12 months; Group 1: mean 1794 mL/min (SD 561); n=26, Group 2: mean 1433 mL/min (SD 521); n=21; Comments: Baseline enalapril: 1690 (561). Baseline placebo: 1437 (521).

Risk of bias: All domain - Very high, Selection - Very high, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: People in the placebo arm appear to have more severe valve disease and worse exercise tolerance than the enalapril arm.; Group 1 Number missing: 6, Reason: 5 withdrew due to nonadherence, 1 developed symptoms and had valve replacement surgery; Group 2 Number missing: 1, Reason: 1 withdrew due to nonadherence

Protocol outcomes not reported by the study	All-cause mortality at ≥12 months; Cardiac mortality at ≥12 months; Quality of life
	at ≥12 months; Quality of life at 6 months; Evidence of HVD progression on imaging
	(worsening of disease severity) at ≥12 months; Withdrawal due to adverse events
	at 6 months; Withdrawal due to adverse events at ≥12 months

Study	Scognamiglio 1990 <sup>145</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=72)
Countries and setting	Conducted in Italy; Setting: Outpatient follow up
Line of therapy	1st line
Duration of study	Intervention + follow up: 1 year
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: Doppler color flow imaging and confirmation by cardiac catheterisation
Stratum	Primary aortic regurgitation
Subgroup analysis within study	Not applicable
Inclusion criteria	People with chronic severe aortic stenosis (grade 3+ to 4+) who were asymptomatic.
Exclusion criteria	Atrial fibrillation, diastolic blood pressure >90mmHg, history of recent development or worsening of the aortic regurgitation (within the preceding 6 months), history of coronary artery disease, mixed aortic stenosis and regurgitation, evidence of additional valvular or congenital heart disease by cardiac catheterisation or echocardiographic and Doppler evaluation, or both, and previous vasodilator or diuretic drug or inotropic therapy (previous therapy with cardioactive drugs).
Recruitment/selection of patients	No additional information stated.
Age, gender and ethnicity	Age - Mean (SD): 35.9 (13.3) years. Gender (M:F): 62:10. Ethnicity: Not stated
Further population details	1. Age: <75 years 2. Disease mechanism for aortic and mitral stenosis: Not applicable 3. Presence vs. absence of uncontrolled systemic hypertension (140/85 mmHg) at the end of trial intervention despite medication: Not stated / Unclear (At the start of the study, nifedipine: 154/60 (19/10), placebo: 155/62 (22/12). No measurements after this. It is likely that the placebo group continued to have hypertension, but no clear reporting of this.).
Indirectness of population	No indirectness
Interventions	(n=38) Intervention 1: Calcium-channel blockers (CCB) - Nifedipine. Nifedipine 20mg twice daily. Duration 1 year. Concurrent medication/care: No cardioactive therapies. No other information provided. Indirectness: No indirectness
	(n=34) Intervention 2: Placebo. Placebo. Duration 1 year. Concurrent medication/care: No cardioactive therapies. No other information provided.

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	Indirectness: No indirectness	
Funding	Funding not stated	
RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: NIFEDIPINE versus PLACEBO  Protocol outcome 1: Need for heart valve intervention at ≥12 months - Actual outcome for Primary aortic regurgitation: Surgery at 1 year; Group 1: 0/36, Group 2: 0/34 Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 2, Reason: 1 withdrew due to adverse events (leg oedema), 1 refused the monthly return visit after 3 months of therapy.; Group 2 Number missing: 0		
Protocol outcome 2: Withdrawal due to adverse events at ≥12 months - Actual outcome for Primary aortic regurgitation: Withdrawal due to adverse event at 1 year; Group 1: 1/38, Group 2: 0/34 Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 0; Group 2 Number missing: 0		
Protocol outcomes not reported by the study	All-cause mortality at ≥12 months; Cardiac mortality at ≥12 months; Quality of life at 6 months; Quality of life at ≥12 months; Onset of symptoms or progression of NYHA class at ≥12 months; Evidence of HVD progression on imaging (worsening of disease severity) at ≥12 months; Exercise tolerance at ≥12 months; Withdrawal due to adverse events at 6 months	

Study	Scognamiglio 1994 <sup>146</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=143)
Countries and setting	Conducted in Italy; Setting: Secondary care
Line of therapy	1st line
Duration of study	Intervention + follow up: 6 years
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: Isolated, chronic, severe aortic regurgitation and normal left ventricular systolic function confirmed by Doppler colour-flow imaging
Stratum	Primary aortic regurgitation
Subgroup analysis within study	Not applicable
Inclusion criteria	Asymptomatic people with isolated, chronic, severe aortic regurgitation and normal left ventricular systolic function.
Exclusion criteria	Recent development or worsening of aortic regurgitation (within the preceding six months); diastolic blood pressure above 90mmHg, a history of coronary artery disease; mixed aortic stenosis and regurgitation (valve gradients ≥20mmHg); evidence of additional valvular or congenital heart disease on echocardiographic or Doppler study; absence of high-quality echocardiographic study of the left ventricle; and an abnormal left ventricular ejection fraction (<50%).
Recruitment/selection of patients	Consecutive people seen at the University of Padua, Italy
Age, gender and ethnicity	Age - Mean (SD): 35.0 (13.0) years. Gender (M:F): 122:21. Ethnicity: Not stated
Further population details	1. Age: <75 years 2. Disease mechanism for aortic and mitral stenosis: Not applicable 3. Presence vs. absence of uncontrolled systemic hypertension (140/85 mmHg) at the end of trial intervention despite medication: Not stated / Unclear (Pretreatment blood pressure - Digoxin: 150/58 (22/14). Nifedipine: 154/60 (20/8).).
Indirectness of population	No indirectness
Interventions	(n=74) Intervention 1: Digoxin. Digoxin 0.25mg daily. Duration 6 years. Concurrent medication/care: No additional information provided. Indirectness: No indirectness (n=69) Intervention 2: Calcium-channel blockers (CCB) - Nifedipine. Nifedipine 20mg twice daily. Duration 6 years. Concurrent medication/care: No additional information. Indirectness: No indirectness

Funding Funding not stated

### RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: DIGOXIN versus NIFEDIPINE

Protocol outcome 1: All-cause mortality at ≥12 months

- Actual outcome for Primary aortic regurgitation: Perioperative death at 6 years; Group 1: 1/70, Group 2: 0/65
Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low,
Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: Reports age, sex, cardiothoracic ratio,
blood pressure and left ventricular echocardiographic parameters; Blinding details: No explanation about blinding. Likely not blinded for caregivers due to
different risks/adverse events from using the different drugs; Group 1 Number missing: 4, Reason: Did not return for the scheduled follow up visits; Group
2 Number missing: 4, Reason: Did not return for the scheduled follow up visits

Protocol outcome 2: Onset of symptoms or progression of NYHA class at ≥12 months

- Actual outcome for Primary aortic regurgitation: Onset of symptoms at 6 years; Group 1: 17/70, Group 2: 6/65
Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low,
Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: Reports age, sex, cardiothoracic ratio,
blood pressure and left ventricular echocardiographic parameters; Blinding details: No explanation about blinding. Likely not blinded for caregivers due to
different risks/adverse events from using the different drugs; Group 1 Number missing: 4, Reason: Did not return for the scheduled follow up visits; Group
2 Number missing: 4, Reason: Did not return for the scheduled follow up visits

Protocol outcome 3: Evidence of HVD progression on imaging (worsening of disease severity) at ≥12 months

- Actual outcome for Primary aortic regurgitation: Left ventricular ejection fraction below 50% at 6 years; Group 1: 5/70, Group 2: 0/65 Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: Reports age, sex, cardiothoracic ratio, blood pressure and left ventricular echocardiographic parameters; Blinding details: No explanation about blinding. Likely not blinded for caregivers due to different risks/adverse events from using the different drugs; Group 1 Number missing: 4, Reason: Did not return for the scheduled follow up visits; Group 2 Number missing: 4, Reason: Did not return for the scheduled follow up visits

Protocol outcome 4: Need for heart valve intervention at ≥12 months

- Actual outcome for Primary aortic regurgitation: Aortic valve replacement at 6 years; Group 1: 20/70, Group 2: 6/65
Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low,
Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Baseline details: Reports age, sex, cardiothoracic ratio,
blood pressure and left ventricular echocardiographic parameters; Blinding details: No explanation about blinding. Likely not blinded for caregivers due to
different risks/adverse events from using the different drugs; Group 1 Number missing: 4, Reason: Did not return for the scheduled follow up visits; Group
2 Number missing: 4, Reason: Did not return for the scheduled follow up visits

Protocol outcome 5: Withdrawal due to adverse events at ≥12 months

- Actual outcome for Primary aortic regurgitation : Withdrawal due to adverse events at 6 years; Group 1: 0/70, Group 2: 0/65

Protocol outcomes not reported by the study

Cardiac mortality at ≥12 months; Quality of life at 6 months; Quality of life at ≥12 months; Exercise tolerance at ≥12 months; Withdrawal due to adverse events at 6 months

Study (subsidiary papers)	SEAS trial: Rossebo 2008 <sup>132</sup> (Bang 2012 <sup>14</sup> , Greve 2019 <sup>59</sup> , Greve 2018 <sup>58</sup> , Greve 2014 <sup>60</sup> , Holme 2010 <sup>71</sup> , Rossebø 2008 <sup>131</sup> )
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=1873)
Countries and setting	Conducted in Denmark, Finland, Germany, Norway, United Kingdom; Setting: Outpatient follow up
Line of therapy	1st line
Duration of study	Intervention + follow up: Median: 52.2 months (4.35 years)
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: Confirmed by echocardiography
Stratum	Primary aortic [including bicuspid] stenosis: While stating it includes mild to moderate aortic stenosis, mean aortic valve area in simvastatin-ezetimibe group = 1.29 (0.48), placebo group = 1.27 (0.46), which are of moderate severity according to British Society of Echocardiography guidance.
Subgroup analysis within study	Not applicable
Inclusion criteria	People between the ages of 45 and 85 years who had asymptomatic, mild-to-moderate aortic valve stenosis, as assessed on echocardiography with a peak aortic-jet velocity of 2.5 to 4m/s.
Exclusion criteria	Previous diagnosis or symptoms of coronary artery disease, peripheral arterial disease, cerebrovascular disease, or diabetes mellitus. If they had any other condition requiring lipid-lowering therapy.
Recruitment/selection of patients	Recruited from five countries across multiple centers.
Age, gender and ethnicity	Age - Mean (SD): 67.6 (9.6). Gender (M:F): 1150:723. Ethnicity: 99.8% of people were white
Further population details	1. Age: Mixed (Based on standard deviation and mean age). 2. Disease mechanism for aortic and mitral stenosis: Mixed (5% had bicuspid aortic valve disease. No statement regarding other aetiology.). 3. Presence vs. absence of uncontrolled systemic hypertension (140/85 mmHg) at the end of trial intervention despite medication: Not stated / Unclear (Blood pressure at the start of the trial. Simvastatin-Ezetimibe arm: 145.6/82.0 (20.4/10.6). Placebo arm: 144.0/82.0 (20.0/10.0). This is in the uncontrolled range. Unclear whether this changes during the trial.).
Indirectness of population	No indirectness

Interventions	(n=944) Intervention 1: Statins - Simvastatin . Simvastatin 40-80mg per day with Ezetimibe 10mg daily. Duration 4.35 years. Concurrent medication/care: Before starting the study, all people were given a single-blind placebo tablet and instructed to follow a lipid-lowering diet. Indirectness: Serious indirectness; Indirectness comment: Includes Ezetimibe 10mg daily combined with a statin (n=929) Intervention 2: Placebo. Placebo. Duration 4.35 years. Concurrent medication/care: Before starting the study, all people were given a single-blind placebo tablet and instructed to follow a lipid-lowering diet. Indirectness: No indirectness
Funding	Study funded by industry (Supported by Merck and Schering-Ploug Pharmaceuticals. Individuals authors supported by a variety of industry bodies.)

### RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: SIMVASTATIN versus PLACEBO

Protocol outcome 1: All-cause mortality at ≥12 months

- Actual outcome for Primary aortic [including bicuspid] stenosis: Death from any cause at 4.35 years; Group 1: 105/944, Group 2: 100/929 Risk of bias: All domain High, Selection High, Blinding Low, Incomplete outcome data Low, Outcome reporting Low, Measurement Low, Crossover Low, Subgroups Low, Other 1 Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 0, Reason: 944 people were assigned to the treatment. 943 received the study drugs. 5 discontinued the study for "other reasons". 198 discontinued study drugs and were followed per protocol. 105 died. All were included in the primary analysis.; Group 2 Number missing: 0, Reason: 929 were assigned to the placebo. All received the placebo. 11 discontinued the study (2 were lost to follow up, 9 had other reasons), 170 discontinued placebo and were followed per protocol, 100 died. All were included in the primary and safety analysis.
- Actual outcome for Primary aortic [including bicuspid] stenosis: Death from any cause at 4.35 years; Group 1: Observed events 105 n=944; Group 2: Observed events 100 n=929; HR 1.04; Lower CI 0.79 to Upper CI 1.36; Log rank variance: 0.80 Risk of bias: All domain High, Selection High, Blinding Low, Incomplete outcome data Low, Outcome reporting Low, Measurement Low, Crossover Low, Subgroups Low, Other 1 Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 0, Reason: 944 people were assigned to the treatment. 943 received the study drugs. 5 discontinued the study for "other reasons". 198 discontinued study drugs and were followed per protocol. 105 died. All were included in the primary analysis.; Group 2 Number missing: 0, Reason: 929 were assigned to the placebo. All received the placebo. 11 discontinued the study (2 were lost to follow up, 9 had other reasons), 170 discontinued placebo and were followed per protocol, 100 died. All were included in the primary analysis.
- Actual outcome for Primary aortic [including bicuspid] stenosis: Death from cardiovascular causes at 4.35 years; Group 1: Observed events 47 n=944; Group 2: Observed events 56 n=929; HR 0.83; Lower CI 0.56 to Upper CI 1.22; Log rank variance: 0.34 Risk of bias: All domain High, Selection High, Blinding Low, Incomplete outcome data Low, Outcome reporting Low, Measurement Low, Crossover Low, Subgroups Low, Other 1 Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 0, Reason: 944 people were assigned to the treatment. 943 received the study drugs. 5 discontinued the study for "other reasons". 198 discontinued study drugs and were followed per

protocol. 105 died. All were included in the primary analysis.; Group 2 Number missing: 0, Reason: 929 were assigned to the placebo. All received the placebo. 11 discontinued the study (2 were lost to follow up, 9 had other reasons), 170 discontinued placebo and were followed per protocol, 100 died. All were included in the primary and safety analysis.

Protocol outcome 2: Cardiac mortality at ≥12 months

- Actual outcome for Primary aortic [including bicuspid] stenosis: Death from cardiovascular causes at 4.35 years; Group 1: 47/944, Group 2: 56/929 Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 0, Reason: 944 people were assigned to the treatment. 943 received the study drugs. 5 discontinued the study for "other reasons". 198 discontinued study drugs and were followed per protocol. 105 died. All were included in the primary analysis.; Group 2 Number missing: 0, Reason: 929 were assigned to the placebo. All received the placebo. 11 discontinued the study (2 were lost to follow up, 9 had other reasons), 170 discontinued placebo and were followed per protocol, 100 died. All were included in the primary and safety analysis.

Protocol outcome 3: Onset of symptoms or progression of NYHA class at ≥12 months

- Actual outcome for Primary aortic [including bicuspid] stenosis: Congestive heart failure as a result of progression of aortic stenosis at 4.35 years; Group 1: 25/944, Group 2: 23/929

Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 0, Reason: 944 people were assigned to the treatment. 943 received the study drugs. 5 discontinued the study for "other reasons". 198 discontinued study drugs and were followed per protocol. 105 died. All were included in the primary analysis.; Group 2 Number missing: 0, Reason: 929 were assigned to the placebo. All received the placebo. 11 discontinued the study (2 were lost to follow up, 9 had other reasons), 170 discontinued placebo and were followed per protocol, 100 died. All were included in the primary and safety analysis.

- Actual outcome for Primary aortic [including bicuspid] stenosis: Congestive heart failure as a result of progression of aortic stenosis at 4.35 years; Group 1: Observed events 25 n=944; Group 2: Observed events 23 n=929; HR 1.09; Lower CI 0.62 to Upper CI 1.92; Log rank variance: 0.77 Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 0, Reason: 944 people were assigned to the treatment. 943 received the study drugs. 5 discontinued the study for "other reasons". 198 discontinued study drugs and were followed per protocol. 105 died. All were included in the primary analysis.; Group 2 Number missing: 0, Reason: 929 were assigned to the placebo. All received the placebo. 11 discontinued the study (2 were lost to follow up, 9 had other reasons), 170 discontinued placebo and were followed per protocol, 100 died. All were included in the primary and safety analysis.

Protocol outcome 4: Need for heart valve intervention at ≥12 months

- Actual outcome for Primary aortic [including bicuspid] stenosis: Aortic valve replacement surgery at 4.35 years; Group 1: 267/944, Group 2: 278/929 Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 0, Reason: 944 people were assigned to the treatment. 943 received the study drugs. 5 discontinued the study for "other reasons". 198 discontinued study drugs and were followed per protocol. 105 died. All were included in the primary analysis.; Group 2 Number missing: 0, Reason: 929 were assigned to the placebo. All received the placebo. 11 discontinued the study (2 were lost to follow up, 9 had other reasons), 170 discontinued placebo and were followed per protocol, 100 died. All

were included in the primary and safety analysis.

- Actual outcome for Primary aortic [including bicuspid] stenosis : Aortic valve replacement surgery at 4.35 years; Group 1: Observed events 267 n=944 ; Group 2: Observed events 278 n=929; HR 1; Lower CI 0.84 to Upper CI 1.18; Log rank variance: 0.97

Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 0, Reason: 944 people were assigned to the treatment. 943 received the study drugs. 5 discontinued the study for "other reasons". 198 discontinued study drugs and were followed per protocol. 105 died. All were included in the primary analysis.; Group 2 Number missing: 0, Reason: 929 were assigned to the placebo. All received the placebo. 11 discontinued the study (2 were lost to follow up, 9 had other reasons), 170 discontinued placebo and were followed per protocol, 100 died. All were included in the primary and safety analysis.

Protocol outcome 5: Withdrawal due to adverse events at ≥12 months

- Actual outcome for Primary aortic [including bicuspid] stenosis: Event resulting in permanent discontinuation of study treatment at 4.35 years; Group 1: 144/943, Group 2: 122/929

Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 1, Reason: 944 people were assigned to the treatment. 943 received the study drugs. 5 discontinued the study for "other reasons". 198 discontinued study drugs and were followed per protocol. 105 died. All but one person was included in the safety analysis (the person who did not receive the medication).; Group 2 Number missing: 0, Reason: 929 were assigned to the placebo. All received the placebo. 11 discontinued the study (2 were lost to follow up, 9 had other reasons), 170 discontinued placebo and were followed per protocol, 100 died. All were included in the primary and safety analysis.

Protocol outcomes not reported by the study

Quality of life at 6 months; Quality of life at ≥12 months; Evidence of HVD progression on imaging (worsening of disease severity) at ≥12 months; Exercise tolerance at ≥12 months; Withdrawal due to adverse events at 6 months

Study	Stewart 2008 <sup>154</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=65)
Countries and setting	Conducted in New Zealand; Setting: Outpatient follow up
Line of therapy	1st line
Duration of study	Intervention + follow up: Median: 19 months (IQR: 15-25 months)
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: Measured by Doppler ultrasound and echocardiography
Stratum	Primary aortic [including bicuspid] stenosis
Subgroup analysis within study	Not applicable
Inclusion criteria	Asymptomatic, moderate or severe aortic stenosis defined as a peak velocity of >3.0 m/s measured by Doppler ultrasound and normal LV systolic function by echocardiography (EF >50%).
Exclusion criteria	Angina, exertional dizziness, syncope, or dyspnoea thought to be related to aortic stenosis, previous or scheduled aortic valve replacement, another heart valve lesion of moderate or greater severity, use of potassium sparing diuretics, serum creatinine level of >0.13 mmol/L or serum potassium level of >5.0 mmol/L during the screening period, significant comorbidity, likely poor compliance, or a contraindication to magnetic resonance imaging.
Recruitment/selection of patients	Not stated
Age, gender and ethnicity	Age - Mean (SD): 67.5 (10.1). Gender (M:F): 50:15. Ethnicity: Not stated
Further population details	1. Age: Mixed (Confidence intervals fall either side of 75 years.). 2. Disease mechanism for aortic and mitral stenosis: Not stated / Unclear 3. Presence vs. absence of uncontrolled systemic hypertension (140/85 mmHg) at the end of trial intervention despite medication: Not stated / Unclear (At the start: Blood pressure eplerenone: 145/83 (21/10). Blood pressure placebo: 144/81 (15/11). Final blood pressure not mentioned.).
Extra comments	
Indirectness of population	No indirectness
Interventions	(n=33) Intervention 1: Diuretics - Eplerenone. Initially epleronone 50mg (1 tablet daily), increased to 100mg (2 tablets daily) after one month if serum potassium level was ≤5.0 mmol/L, serum creatinine level was ≤0.13 mmol/L, and systolic blood pressure was >100mmHg, and there were no adverse events of treatment. Duration

	19 months. Concurrent medication/care: Other medications were at the discretions of the patient's usual doctor. Indirectness: No indirectness  (n=32) Intervention 2: Placebo. Placebo. Duration 19 months. Concurrent medication/care: Other medications were at the discretions of the patient's usual doctor. Indirectness: No indirectness
Funding	Study funded by industry (The National Heart Foundation of New Zealand and Pfizer (states it was completed by the study investigators independently from sponsors). The Green Lane Research and Education fund used for salary support)

#### RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: EPLERENONE versus PLACEBO

Protocol outcome 1: All-cause mortality at ≥12 months

- Actual outcome for Primary aortic [including bicuspid] stenosis : All-cause mortality at 19 months; Group 1: 1/30, Group 2: 2/31
- Risk of bias: All domain Low, Selection Low, Blinding Low, Incomplete outcome data Low, Outcome reporting Low, Measurement Low, Crossover
- Low, Subgroups Low, Other 1 Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 2, Reason: 2 withdrawals due to patient decision, 1 withdrawal due to gynaecomastia; Group 2 Number missing: 1, Reason: 1 withdrawal due to patient decision

### Protocol outcome 2: Cardiac mortality at ≥12 months

- Actual outcome for Primary aortic [including bicuspid] stenosis : Sudden death at 19 months; Group 1: 0/29, Group 2: 1/30
- Risk of bias: All domain Low, Selection Low, Blinding Low, Incomplete outcome data Low, Outcome reporting Low, Measurement Low, Crossover Low, Subgroups Low, Other 1 Low, Indirectness of outcome: No indirectness: Group 1 Number missing: 2 Reason: 2 withdrawals due to patient
- Low, Subgroups Low, Other 1 Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 2, Reason: 2 withdrawals due to patient decision, 1 withdrawal due to gynaecomastia; Group 2 Number missing: 1, Reason: 1 withdrawal due to patient decision

# Protocol outcome 3: Quality of life at ≥12 months

- Actual outcome for Primary aortic [including bicuspid] stenosis: SF36 physical functioning subscale at 19 months; Group 1: mean -5 (SD 22); n=29, Group 2: mean -9 (SD 19); n=30; SF-36 physical functioning subscale 0-100 Top=High is good outcome; Comments: Baseline eplerenone: 79 (22). Baseline placebo: 87 (10).
- Risk of bias: All domain Low, Selection Low, Blinding Low, Incomplete outcome data Low, Outcome reporting Low, Measurement Low, Crossover Low, Subgroups Low, Other 1 Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 4, Reason: 2 withdrawals due to patient decision, 1 withdrawal due to gynaecomastia, 1 non-cardiac death; Group 2 Number missing: 2, Reason: 1 withdrawal due to patient decision, 1 non-cardiac death
- Actual outcome for Primary aortic [including bicuspid] stenosis: SF36 role physical subscale at 19 months; Group 1: mean -9 (SD 34); n=29, Group 2: mean -12 (SD 37); n=30; SF-36 role physical subscale 0-100 Top=High is good outcome; Comments: Baseline eplerenone: 72 (42). Baseline placebo: 82 (30).
- Risk of bias: All domain Low, Selection Low, Blinding Low, Incomplete outcome data Low, Outcome reporting Low, Measurement Low, Crossover Low, Subgroups Low, Other 1 Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 4, Reason: 2 withdrawals due to patient

decision, 1 withdrawal due to gynaecomastia, 1 non-cardiac death; Group 2 Number missing: 2, Reason: 1 withdrawal due to patient decision, 1 non-cardiac death

Protocol outcome 4: Onset of symptoms or progression of NYHA class at ≥12 months

- Actual outcome for Primary aortic [including bicuspid] stenosis: Symptomatic deterioration at 19 months; Group 1: 13/29, Group 2: 10/30 Risk of bias: All domain Low, Selection Low, Blinding Low, Incomplete outcome data Low, Outcome reporting Low, Measurement Low, Crossover Low, Subgroups Low, Other 1 Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 4, Reason: 2 withdrawals due to patient
- Low, Subgroups Low, Other 1 Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 4, Reason: 2 withdrawals due to patient decision, 1 withdrawal due to gynaecomastia, 1 non-cardiac death; Group 2 Number missing: 2, Reason: 1 withdrawal due to patient decision, 1 non-cardiac death

Protocol outcome 5: Withdrawal due to adverse events at ≥12 months

- Actual outcome for Primary aortic [including bicuspid] stenosis: Withdrawal due to adverse events at 19 months; Group 1: 1/32, Group 2: 0/30 Risk of bias: All domain Low, Selection Low, Blinding Low, Incomplete outcome data Low, Outcome reporting Low, Measurement Low, Crossover
- Low, Subgroups Low, Other 1 Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 3, Reason: 2 withdrawals due to patient decision, 1 non-cardiac death; Group 2 Number missing: 2, Reason: 1 withdrawal due to patient decision, 1 non-cardiac death

Protocol outcomes not reported by the study

Quality of life at 6 months; Evidence of HVD progression on imaging (worsening of disease severity) at ≥12 months; Need for heart valve intervention at ≥12 months; Exercise tolerance at ≥12 months; Withdrawal due to adverse events at 6 months

Study	TASS trial: Dichtl 2008 <sup>41</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=50)
Countries and setting	Conducted in Austria; Setting: Outpatient care
Line of therapy	1st line
Duration of study	Intervention + follow up: 3-5 years
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: Transthoracic echocardiography completed throughout the study
Stratum	Primary aortic [including bicuspid] stenosis
Subgroup analysis within study	Not applicable
Inclusion criteria	People aged >18 years with calcific aortic stenosis, mean systolic gradients of ≥18 mmHg, valvular stenotic flow velocities ≥20m/s and aortic valve calcification on echo.
Exclusion criteria	Child-bearing potential, severe liver disease, concomitant mitral valve stenosis, severe liver disease, concomitant mitral valve stenosis, severe mitral or aortic regurgitation, advanced left ventricular dysfunction (ejection fraction <40%), planned aortic valve replacement, intolerance of statins, or an indication for statin therapy according to guidelines.
Recruitment/selection of patients	Consecutively referred 120 people to their echocardiographic laboratory for evaluation of asymptomatic calcified aortic stenosis. 50 were enrolled and followed up every 12 months for 3-5 years.
Age, gender and ethnicity	Age - Mean (SD): 67.0 (11.7) years. Gender (M:F): 28:22. Ethnicity: Not stated
Further population details	1. Age: Mixed (Confidence interval falls over 75 years. Intervention: 64.2 (12.0), Control: 69.7 (10.6)). 2. Disease mechanism for aortic and mitral stenosis: Calcific 3. Presence vs. absence of uncontrolled systemic hypertension (140/85 mmHg) at the end of trial intervention despite medication: Not stated / Unclear (States that 9 in the intervention arm, and 14 in the control arm had hypertension. Antihypertensive medication was prescribed. Does not state if this was uncontrolled at the end of the trial.).
Indirectness of population	No indirectness
Interventions	(n=25) Intervention 1: Statins - Atorvastatin . Atorvastatin 20mg once a day. Duration 3-5 years. Concurrent medication/care: 7 people were using aspirin, 6 people were using an ACE inhibitor, 1 person was using a calcium channel blocker,

(n=25) Intervention 2: Placebo. Placebo. Duration 3-5 years. Concurrent medication/care: 14 people were using aspirin, 11 were using an ACE inhibitor, 2 were using a calcium channel blocker, 5 were using a beta blocker, 3 were using a vitamin K antagonist. Indirectness: No indirectness		1 person with using a beta blocker, 1 person was using a vitamin K antagonist. Indirectness: No indirectness
Funding Faultment / drugs provided by industry (Medication provided by Pfizer Austria)		medication/care: 14 people were using aspirin, 11 were using an ACE inhibitor, 2 were using a calcium channel blocker, 5 were using a beta blocker, 3 were using a
Equipment 7 drugs provided by industry (inedication provided by Frizer Austria)	Funding	Equipment / drugs provided by industry (Medication provided by Pfizer Austria)

#### RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: ATORVASTATIN versus PLACEBO

Protocol outcome 1: All-cause mortality at ≥12 months

- Actual outcome for Primary aortic [including bicuspid] stenosis: All-cause mortality at 5 years; Group 1: 1/24, Group 2: 1/23
Risk of bias: All domain - Very high, Selection - Very high, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low; Indirectness of outcome: No indirectness; Baseline details: Reports age, gender, coronary artery calcification, arterial hypertension, current smoking, renal insufficiency and medication use. Statin arm has people less medication usage, but more coronary artery disease and smoking, control arm has more people with arterial hypertension.; Group 1 Number missing: 1, Reason: 1 lost due to an intolerance to atorvastatin therapy; Group 2 Number missing: 2, Reason: 1 lost to follow up, 1 developed gastric cancer and left the trial

# Protocol outcome 2: Cardiac mortality at ≥12 months

- Actual outcome for Primary aortic [including bicuspid] stenosis: Cardiac mortality at 5 years; Group 1: 1/24, Group 2: 1/23
Risk of bias: All domain - Very high, Selection - Very high, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low; Indirectness of outcome: No indirectness; Baseline details: Reports age, gender, coronary artery calcification, arterial hypertension, current smoking, renal insufficiency and medication use. Statin arm has people less medication usage, but more coronary artery disease and smoking, control arm has more people with arterial hypertension.; Group 1 Number missing: 1, Reason: 1 lost due to an intolerance to atorvastatin therapy; Group 2 Number missing: 2, Reason: 1 lost to follow up, 1 developed gastric cancer and left the trial

#### Protocol outcome 3: Need for heart valve intervention at ≥12 months

- Actual outcome for Primary aortic [including bicuspid] stenosis: Need for heart valve intervention at 5 years; Group 1: 5/24, Group 2: 1/23 Risk of bias: All domain - Very high, Selection - Very high, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low; Indirectness of outcome: No indirectness; Baseline details: Reports age, gender, coronary artery calcification, arterial hypertension, current smoking, renal insufficiency and medication use. Statin arm has people less medication usage, but more coronary artery disease and smoking, control arm has more people with arterial hypertension.; Group 1 Number missing: 1, Reason: 1 lost due to an intolerance to atorvastatin therapy; Group 2 Number missing: 2, Reason: 1 lost to follow up, 1 developed gastric cancer and left the trial

#### Protocol outcome 4: Withdrawal due to adverse events at 6 months

- Actual outcome for Primary aortic [including bicuspid] stenosis: Withdrawal due to adverse events at <6 months; Group 1: 1/25, Group 2: 0/23

Risk of bias: All domain - Very high, Selection - Very high, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low; Indirectness of outcome: No indirectness; Baseline details: Reports age, gender, coronary artery calcification, arterial hypertension, current smoking, renal insufficiency and medication use. Statin arm has people less medication usage, but more coronary artery disease and smoking, control arm has more people with arterial hypertension.; Group 1 Number missing: 0, Reason: 1 lost due to an intolerance to atorvastatin therapy; Group 2 Number missing: 2, Reason: 1 lost to follow up, 1 developed gastric cancer and left the trial

Protocol outcomes not reported by the study

Quality of life at 6 months; Quality of life at ≥12 months; Onset of symptoms or progression of NYHA class at ≥12 months; Evidence of HVD progression on imaging (worsening of disease severity) at ≥12 months; Exercise tolerance at ≥12 months; Withdrawal due to adverse events at ≥12 months

Study	Wisenbaugh 1994-1 <sup>177</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=32)
Countries and setting	Conducted in South Africa; Setting: Outpatient follow up
Line of therapy	1st line
Duration of study	Intervention + follow up: 6 months
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: Doppler echocardiography and clinical examination
Stratum	Primary mitral regurgitation
Subgroup analysis within study	Not applicable
Inclusion criteria	Severe isolated MR by clinical examination and Doppler echocardiographic criteria; mitral valve are >3cm² and no aortic stenosis or other significant valvular lesion; normal sinus rhythm; no clinical evidence of coronary artery disease; clear endocardial borders identifiable on echocardiographic imaging; willingness to participate in the protocol and high probability of good follow up as determined by a nurse who interviewed the person in their own language.
Exclusion criteria	More than mild symptoms (>NYHA class II).
Recruitment/selection of patients	People recruited from the cardiac clinic at Baragwanath Hospital
Age, gender and ethnicity	Age - Other: Mean age: 24.9. Gender (M:F): 5:25. Ethnicity:
Further population details	1. Age: <75 years (Mean age captopril: 26, mean age placebo: 24). 2. Disease mechanism for aortic and mitral stenosis: Not applicable 3. Presence vs. absence of uncontrolled systemic hypertension (140/85 mmHg) at the end of trial intervention despite medication: Not stated / Unclear (initial blood pressure captopril: 117/67, placebo: 110/63. From the graph it appears the value stayed under 140/85 but it is unclear.).
Extra comments	Three had myxomatous MR, the rest had rheumatic MR.
Indirectness of population	No indirectness
Interventions	(n=14) Intervention 1: Angiotensin-converting enzyme (ACE) inhibitors - Captopril. Captopril 25mg three times daily. Duration 6 months. Concurrent medication/care: Three had been taking enalapril (5-10mg twice daily), which was discontinued at least 2 months before entry into the study. One was taking nifedipine which was discontinued 1 month prior to entry. Other vasodilating and digitalis drugs were not used. People who were on furosemide were maintained on a constant dose

	(average captopril group: 46mg, average placebo group: 42mg). Indirectness: No indirectness  (n=18) Intervention 2: Placebo. Placebo. Duration 6 months. Concurrent medication/care: Three had been taking enalapril (5-10mg twice daily), which was discontinued at least 2 months before entry into the study. One was taking nifedipine which was discontinued 1 month prior to entry. Other vasodilating and digitalis drugs were not used. People who were on furosemide were maintained on a constant dose (average captopril group: 46mg, average placebo group: 42mg). Indirectness: No indirectness
Funding	Funding not stated

### RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: CAPTOPRIL versus PLACEBO

Protocol outcome 1: All-cause mortality at ≥12 months

- Actual outcome for Primary mitral regurgitation: Death at 6 months; Group 1: 0/12, Group 2: 1/17
Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low,
Crossover - Low, Subgroups - Low, Other 1 - Low, Comments - Reports that there was a randomisation error; Indirectness of outcome: No indirectness;
Group 1 Number missing: 2, Reason: 2 excluded due to poor compliance; Group 2 Number missing: 1, Reason: 1 excluded due to poor compliance

Protocol outcome 2: Cardiac mortality at ≥12 months

- Actual outcome for Primary mitral regurgitation: Death at 6 months; Group 1: 0/12, Group 2: 1/17

Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low, Comments - Reports that there was a randomisation error; Indirectness of outcome: No indirectness; Group 1 Number missing: 2, Reason: 2 excluded due to poor compliance; Group 2 Number missing: 1, Reason: 1 excluded due to poor compliance

Protocol outcome 3: Onset of symptoms or progression of NYHA class at ≥12 months

- Actual outcome for Primary mitral regurgitation: Symptom deterioration at 6 months; Group 1: 0/12, Group 2: 1/17
Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Other 1 - Low, Comments - Reports that there was a randomisation error; Indirectness of outcome: No indirectness; Group 1 Number missing: 2, Reason: 2 excluded due to poor compliance; Group 2 Number missing: 1, Reason: 1 excluded due to poor compliance

Quality of life at 6 months; Quality of life at ≥12 months; Evidence of HVD
progression on imaging (worsening of disease severity) at ≥12 months; Need for
heart valve intervention at ≥12 months; Exercise tolerance at ≥12 months;

Withdrawal due to adverse events at 6 months; Withdrawal due to adverse events at ≥12 months

Study	Wisenbaugh 1994-2 <sup>177</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=23)
Countries and setting	Conducted in South Africa; Setting: Outpatient follow up
Line of therapy	1st line
Duration of study	Intervention + follow up: 6 months
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: Doppler echocardiography and clinical examination
Stratum	Primary aortic regurgitation
Subgroup analysis within study	Not applicable
Inclusion criteria	Severe isolated AR by clinical examination and Doppler echocardiographic criteria; mitral valve are >3cm² and no aortic stenosis or other significant valvular lesion; normal sinus rhythm; no clinical evidence of coronary artery disease; clear endocardial borders identifiable on echocardiographic imaging; willingness to participate in the protocol and high probability of good follow up as determined by a nurse who interviewed the person in their own language.
Exclusion criteria	More than mild symptoms (>NYHA class II)
Recruitment/selection of patients	People recruited from the cardiac clinic at Baragwanath Hospital
Age, gender and ethnicity	Age - Other: Mean age: 28.1 years. Gender (M:F): 15:5. Ethnicity: Not stated
Further population details	1. Age: <75 years (Mean age Captopril: 29, Placebo: 27). 2. Disease mechanism for aortic and mitral stenosis: Not applicable 3. Presence vs. absence of uncontrolled systemic hypertension (140/85 mmHg) at the end of trial intervention despite medication: Not stated / Unclear (Baseline captopril: 131/46, Baseline placebo: 144/57. So potentially. According to the graph there wasn't much charge at 6 months, but unclear.).
Extra comments	In most cases underlying cause of AR could not be determined (none were thought to have a rheumatic aetiology).
Indirectness of population	No indirectness

Interventions	(n=13) Intervention 1: Angiotensin-converting enzyme (ACE) inhibitors - Captopril. Captopril 25mg three times a day. Duration 6 months. Concurrent medication/care: In the placebo group, one person had been taken enalapril, which was discontinued for two months before entry. One was taking hydralazine which was discontinued for one month. In the captopril group, two had been enalapril which was discontinued for two and three months respectively. Three were taking hydralazine, which was discontinued for 2-3 months before entry. People who were on chronic furosemide were maintained at a constant dose (captopril group average daily dose = 47mg, placebo group average daily dose = 31mg)  (n=10) Intervention 2: Placebo. Placebo. Duration 6 months. Concurrent medication/care: In the placebo group, one person had been taken enalapril, which was discontinued for two months before entry. One was taking hydralazine which was discontinued for one month. In the captopril group, two had been enalapril which was discontinued for two and three months respectively. Three were taking hydralazine, which was discontinued for 2-3 months before entry. People who were on chronic furosemide were maintained at a constant dose (captopril group average daily dose = 47mg, placebo group average daily dose = 31mg). Indirectness: No indirectness
Funding	Funding not stated

# RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: CAPTOPRIL versus PLACEBO

Protocol outcome 1: Onset of symptoms or progression of NYHA class at ≥12 months

- Actual outcome for Primary aortic regurgitation: Symptom deterioration at 6 months; Group 1: 0/11, Group 2: 0/9
Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low,
Crossover - Low, Subgroups - Low, Other 1 - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 2, Reason: 2 lost to follow up after 3 months; Group 2 Number missing: 1, Reason: 1 lost to follow up after 3 months

# Protocol outcomes not reported by the study

All-cause mortality at ≥12 months; Cardiac mortality at ≥12 months; Quality of life at 6 months; Quality of life at ≥12 months; Evidence of HVD progression on imaging (worsening of disease severity) at ≥12 months; Need for heart valve intervention at ≥12 months; Exercise tolerance at ≥12 months; Withdrawal due to adverse events at 6 months; Withdrawal due to adverse events at ≥12 months

Study	Alan 2002 <sup>6</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	N/A (n=80)
Countries and setting	Conducted in Turkey; Setting: Initiated and followed up in secondary care
Line of therapy	1st line
Duration of study	Intervention + follow up: 3 months (received an IV dose of diltiazem for induction, then 3 months of oral diltiazem)
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: Noted previous diagnosis of mitral stenosis as assessment method for HVD (However, go on to do echocardiography confirming this). Congestive heart failure assessed by NYHA status.
Stratum	Primary mitral stenosis: Noted previous diagnosis of mitral stenosis as assessment method for HVD (However, go on to do echocardiography confirming this).
Subgroup analysis within study	Not applicable: N/A
Inclusion criteria	People with a diagnosis of mitral stenosis of mild-to-moderate severity

Exclusion criteria	People with moderate-to-severe degrees of aortic insufficiency and aortic stenosis, and patients with severe pulmonary hypertension or right-sided heart failure.
Recruitment/selection of patients	No additional information available.
Age, gender and ethnicity	Age - Mean (SD): 38±6.8 years. Gender (M:F): 28:52 (35/65%). Ethnicity: Not stated
Further population details	1. Age: <75 years (Range: 33-45 years). 2. Heart rate: Normal (82±10 per minute). 3. Presence vs. absence of uncontrolled systemic hypertension: Absence of uncontrolled systemic hypertension (Mean 112/71 and 115/78 in diltiazem and metoprolol groups). 4. Severe vs non-severe HVD: Non-severe (Mild-to-moderate). 5. Symptomatic vs asymptomatic: Symptomatic (NYHA II or III).
Extra comments	Age range = 33-45 years; Functional capacities of all patients included in the study were NYHA class II and class III.
Indirectness of population	No indirectness: Population fits guideline condition (adults aged 18 years and over diagnosed with mitral stenosis with congestive heart failure).
Interventions	(n=40) Intervention 1: Beta blockers - Metoprolol. Initially: 5mg intravenous, followed by 50mg orally twice daily. Duration 3 months. Concurrent medication/care: Not stated. Indirectness: No indirectness Comments: Initial intravenous loading dose. Not necessarily standard treatment.

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	(n=40) Intervention 2: Calcium-channel blockers (CCB) - Diltiazem. Initially 25mg intravenously followed by 60mg orally three times daily. Duration 3 months. Concurrent medication/care: Not stated. Indirectness: No indirectness
Funding	Funding not stated

# RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: METOPROLOL versus DILTIAZEM

Protocol outcome 1: Exercise tolerance at 12 months

- Actual outcome for Primary mitral stenosis: Total effort time (sec) - treadmill exercise test (Bruce protocol) at 3 months; Group 1: mean 520 Seconds (SD 90); n=40, Group 2: mean 570 Seconds (SD 126); n=40; Comments: Initial values (at beginning of treatment):

Metoprolol: 452±120 Diltiazem: 534±120

Note: Results are from 3 months, outside of that required for inclusion in the outcome.

Risk of bias: All domain - Very high, Selection - Very high, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Comments - Outcome follow up is at 3 months rather than the 6 or greater required for inclusion.; Indirectness of outcome: No indirectness, Comments: Bruce protocol is widely used in clinical practice.; Baseline details: Difference in baseline effort time greater than difference in final values. Full baseline characteristics not reported. The age and sex is reported, and appears similar.; Group 1 Number missing: 0; Group 2 Number missing: 0

Protocol outcome 2: Withdrawal due to adverse events at 6 months

- Actual outcome for Primary mitral stenosis: Adverse events that interrupted treatment at 3 months; Group 1: 0/40, Group 2: 0/40; Comments: Note: Outcome assessed at 3 months. Outside of the value accepted for follow up in this protocol.

Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low, Comments - Outcome follow up is at 3 months rather than the 6 or greater required for inclusion.; Indirectness of outcome: No indirectness; Baseline details: Full baseline characteristics not reported. The age and sex is reported, and appears similar.; Group 1 Number missing: 0; Group 2 Number missing: 0

Protocol outcomes not reported by the study

Quality of life at 6 months; Hospitalisation due to heart failure at 12 months; Cardiac mortality at 12 months; Quality of life at 12 months; Need for valve intervention at 6 months; Need for valve intervention at 12 months; Withdrawal due to adverse events at 12 months; All-cause mortality at 12 months; Cardiac mortality at 6 months

Study	Bassan 1987 <sup>15</sup>
Study type	RCT (Patient randomised; Crossover: 1 week (washout period), except for week 3 (when patients were given half doses of propranolol - during this period patients could have had propranolol or placebo during the week before with no washout))
Number of studies (number of participants)	N/A (n=10)
Countries and setting	Conducted in Israel; Setting: Not stated - likely secondary care
Line of therapy	1st line
Duration of study	Intervention + follow up: 3 weeks, follow up for at least 24 months after the intervention
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: Well explained. Diagnosis established by typical auscultatory, electrocardiographic, radiographic, and echocardiographic findings.
Stratum	Primary mitral stenosis: Patients with isolated mitral stenosis.
Subgroup analysis within study	Not applicable
Inclusion criteria	Patients with isolated mitral stenosis.
Exclusion criteria	No patient had echocardiographic evidence of left ventricular hypertrophy or abnormal left ventricular function. No evidence of right heart failure. No obstructive lung disease or contraindication to beta-blockade.
Recruitment/selection of patients	No information stated
Age, gender and ethnicity	Age - Other: Individual values given. The mean is 38.7 (Range: 19-56). Gender (M:F): 4:6. Ethnicity: Not stated
Further population details	1. Age: <75 years (Mean = 38.7). 2. Heart rate: Not stated / Unclear (No pre-treatment values given; comparison between placebo and treatment group shows bradycardiac-normal rates). 3. Presence vs. absence of uncontrolled systemic hypertension: Not stated / Unclear 4. Severe vs non-severe HVD: Not stated / Unclear (At least 2 with "severe stenosis". Others not clearly stated.). 5. Symptomatic vs asymptomatic: Symptomatic (NYHA class II or III.).
Extra comments	NYHA class II or III. They determine the mitral valve area post-hoc for some cases (after surgery).
Indirectness of population	No indirectness: All patients are older than 18 years. The range is fairly last, but not indirect.
Interventions	(n=10) Intervention 1: Beta blockers - Propranolol . Full dose (dependent on weight) - 40mg orally (two or three times a day dependent on weight). Patients were trained according to an individual exercise protocol to reach a reproducible degree of near maximal dyspnoea. Five patients performed bicycle exercise and five exercised on a treadmill. In the training phase the patients learnt to recognise their dyspnoea end point. Achievement of stable performance (less than a 30 second variation in exercise duration) usually required 8-12 exercise bouts over several weeks with 2-3 bouts per session. The starting level for the study phase was chosen so that endpoint dyspnoea was reached after 3-6 minutes of exercise. At the final training session the patient was given a test dose of

0	
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Protocol outcomes not reported by the

study

	40mg of propranolol and was observed for several hours for possible adverse effects as well as for the degree of induced bradycardia. Duration 1 week. Concurrent medication/care: None stated. Indirectness: No indirectness; Indirectness comment: Is only for a short duration, but is not indirect in itself.
	(n=10) Intervention 2: Placebo. Matching placebo. Duration 1 week. Concurrent medication/care: None stated. Indirectness: No indirectness; Indirectness comment: Is only for a short duration, but is not indirect in itself.
Funding	Academic or government funding (MacRamer Heart Research Scholarship Fund, Flushing, New York)
·	SED) AND RISK OF BIAS FOR COMPARISON: PROPRANOLOL (FULL DOSE) versus PLACEBO
	lerance at 12 months trail stenosis: Time to near maximal dyspnoea at 1 week; Group 1: mean 274 seconds (SD 79); n=10, Group 2: mean 283 ents: SD calculated from SE in paper
	igh, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Low; Indirectness of outcome: No indirectness; Baseline details: Crossover study; Group 1 Number missing: ; Group 2

All-cause mortality at 12 months; Cardiac mortality at 6 months

Quality of life at 6 months; Hospitalisation due to heart failure at 12 months; Cardiac mortality at 12 months;

months; Withdrawal due to adverse events at 6 months; Withdrawal due to adverse events at 12 months;

Quality of life at 12 months; Need for valve intervention at 6 months; Need for valve intervention at 12

Study	Chockalingam 2004 <sup>29</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=56)
Countries and setting	Conducted in India; Setting: Secondary care
Line of therapy	1st line
Duration of study	Intervention + follow up: 12 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: Referral from a cardiology service and states specific parameters that needed to be met for inclusion.
Stratum	Primary aortic [including bicuspid] stenosis: Unclear how defined/diagnosed
Subgroup analysis within study	Post-hoc subgroup analysis: They reported completing subgroup analysis for age, sex, baseline walk distance and LV dysfunction for the outcome of effort tolerance - they showed no difference. They found that patients with associated regurgitant lesions had a trend toward more improvement in exercise capacity and symptoms in another analysis.
Inclusion criteria	Severe AS (aortic valve area <0.75 cm2, mean aortic gradient >50 mm Hg, or aortic valve Doppler jet >4.5 m/s) and symptomatic New York Heart Association class III or IV dyspnoea or angina.
Exclusion criteria	Persistent hypotension (systolic BP <90 or mean BP <60), severe mitral stenosis (mitral valve orifice <1.0 cm2), known intolerance for ACEI, and renal dysfunction (serum creatinine >2.5 mg/dL).
Recruitment/selection of patients	Recruitment from a cardiology service of a medical college hospital.
Age, gender and ethnicity	Age - Mean (SD): Intervention arm: 43±11, Control arm: 46±12. Gender (M:F): 39:13. Ethnicity: Not reported
Further population details	1. Age: <75 years (Mean in Enalapril arm = 43 +/-11, Mean in Placebo arm = 46 +/-12). 2. Heart rate: Normal (Mean in Enalapril arm = 83 +/-8, Mean in Placebo arm = 83 +/-8). 3. Presence vs. absence of uncontrolled systemic hypertension: Absence of uncontrolled systemic hypertension (mean blood pressure for both arms given. Doesn't state whether this is systolic or diastolic. However, for both arms this is less than or equal to 90mmHg). 4. Severe vs non-severe HVD: Severe 5. Symptomatic vs asymptomatic: Symptomatic
Extra comments	Patients were waiting for surgery or unwilling to have surgery.
Indirectness of population	No indirectness: Fits our guideline criteria.
Interventions	(n=37) Intervention 1: Angiotensin-converting enzyme (ACE) inhibitors - Enalapril. 2.5mg BD gradually titrated up to 10mg BD over 2 weeks (in 5 patients this was not achievable and they remained at 2.5-5mg BD during the study).  All were initially stabilised in hospital (5 ± 3 days) with diuretics, digoxin, and intravenous dobutamine infusion before initiating the study medication. Duration 3 months (in the majority). Concurrent

	medication/care: Prior treatment was continued (apart from potassium replacement for which the dose was decreased). All had frusemide, 94% had digoxin, 48% had spironalactone and 9.6% had dobutamine. Indirectness: No indirectness:  (n=19) Intervention 2: Placebo. All were initially stabilised in hospital (5 ± 3 days) with diuretics, digoxin, and intravenous dobutamine infusion before initiating the study medication. Duration 3 months. Concurrent medication/care: Prior treatment was continued. All had frusemide, 94% had digoxin, 48% had spironalactone and 9.6% had dobutamine. Indirectness: No indirectness
Funding	Funding not stated
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RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: ENALAPRIL versus PLACEBO

Protocol outcome 1: Exercise tolerance at 12 months

- Actual outcome for Primary aortic [including bicuspid] stenosis: 6-minute walk distance (meters) at 4 weeks; Group 1: mean 402 Meters (SD 150); n=34, Group 2: mean 376 Meters (SD 174); n=18; Comments: Values at baseline were reported:

Enalapril arm: 330±157 Control arm: 349±147

Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low; Indirectness of outcome: No indirectness; Baseline details: 2:1 ratio of intervention to placebo. However, proportions were maintained.; Group 1 Number missing: 3, Reason: Patients withdrew due to adverse events; Group 2 Number missing: 1, Reason: Patient withdrew due to adverse events

Protocol outcome 3: Withdrawal due to adverse events at 6 months

- Actual outcome for Primary aortic [including bicuspid] stenosis: Withdrawal due to intolerance to study medication at 3 months; Group 1: 3/37, Group 2: 1/19; Comments: Enalapril: 3 had significant hypotension; placebo: unclear

Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low, Subgroups - Low; Indirectness of outcome: No indirectness; Baseline details: 2:1 ratio of intervention to placebo. However, proportions were maintained.; Group 1 Number missing: 0; Group 2 Number missing: 0

Protocol outcomes not reported by the study

Quality of life at 6 and 12 months; Hospitalisation due to heart failure at 12 months; Cardiac mortality at 6 and 12 months; Need for valve intervention at 6 or 12 months; Withdrawal due to adverse events at 12 months; All-cause mortality at 12 months.

Study	Dalsgaard 2014 <sup>38</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=44)

Countries and setting	Conducted in Denmark; Setting: Secondary care
Line of therapy	1st line
Duration of study	Follow up (post intervention): Median of 8 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: Aortic valve area <1cm², in sinus rhythm and without symptoms at rest = severe AS. Independently assessed for NYHA class.
Stratum	Primary aortic [including bicuspid] stenosis
Subgroup analysis within study	Not applicable
Inclusion criteria	Severe AS. Symptomatic and asymptomatic patients (32 were symptomatic, 12 were asymptomatic).
Exclusion criteria	Mitral regurgitation, unable to perform exercise testing, resting systolic BP <100 mmHg, known renal artery stenosis or creatinine >200 umol/l, prior treatment with ACE-I or ARBs in the last month.
Recruitment/selection of patients	Consecutive
Age, gender and ethnicity	Age - Mean (SD): 69.9±8.3 (range of 55-85 years). Gender (M:F): 4:7. Ethnicity: Not stated
Further population details	1. Age: Mixed 2. Heart rate: Not stated / Unclear 3. Presence vs. absence of uncontrolled systemic hypertension: Not stated / Unclear 4. Severe vs non-severe HVD Severe 5. Symptomatic vs asymptomatic: Mixed
Extra comments	Severe aortic stenosis - 32 patients were symptomatic, 12 were asymptomatic. 30 had another comorbidity (hypertension, IHD, diabetes mellitus).
Indirectness of population	No indirectness
Interventions	(n=22) Intervention 1: Angiotensin-converting enzyme (ACE) inhibitors - Trandolapril. Daily increasing doses of trandolapril, 0.5 mg on day 1; 1 mg on day 2 and 2 mg in day 3. At discharge on day 3 patients were given the maximum tolerated dose for the rest of the study. Duration 8 weeks. Concurrent medication/care: Calcium antagonists (14%); beta-blockers (27%); diuretics (50%). Indirectness: No indirectness
	(n=22) Intervention 2: Placebo. Matched placebo. Duration 8 weeks. Concurrent medication/care: Calcium antagonists (18%); beta-blockers (45%); diuretics (32%). Indirectness: No indirectness
Funding	Funding not stated

- Actual outcome for Primary aortic [including bicuspid] stenosis: Exercise duration to exhaustion (minutes) at 3 days; Group 1: mean 0.2 minutes (SD 0.6); n=21, Group 2: mean 0.2 minutes (SD 0.4); n=22; Comments: Baseline: ACE-I: 5.9 (1.9); placebo: 6.1 (2.1) minutes
Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low,
Crossover - Low; Indirectness of outcome: Serious indirectness, Comments: Follow-up period 3 days; Group 1 Number missing: 1, Reason: Ischaemic stroke; Group 2 Number missing: 0

Protocol outcome 3: Withdrawal due to adverse events at 6 months

- Actual outcome for Primary aortic [including bicuspid] stenosis: Withdrawal due to adverse events at 8 weeks; Group 1: 1/22, Group 2: 0/22; Comments: 1 cerebral ischaemic stroke (not thought to be due to the study drug)

Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: Serious indirectness, Comments: Follow-up period 8 weeks; Group 1 Number missing: 0; Group 2 Number missing: 0

Protocol outcomes not reported by the study

Quality of life at 6 months; Hospitalisation due to heart failure at 12 months; Cardiac mortality at 12 months; Quality of life at 12 months; Need for valve intervention at 6 or 12 months; Withdrawal due to adverse events at 12 months; All-cause mortality at 12 months; Cardiac mortality at 6 months

Study	Helske-Suihko 2015 <sup>67</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=51)
Countries and setting	Conducted in Finland; Setting: Secondary care
Line of therapy	1st line
Duration of study	Intervention + follow up: 46 to 310 days. Mean = $164 + /-67$ days for intervention arm, $151 + /-72$ days for control arm (a little under 6 months).
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: Previous diagnosis of AS (have been referred to them for consideration for surgery). However, they have an echo as a part of the intervention, which would assess this. The paper states the majority of patients had symptoms equivalent to NYHA class II, but selectively reports proportion in classes I-II vs. III. No clear assessment of NYHA reported.
Stratum	Primary aortic [including bicuspid] stenosis
Subgroup analysis within study	Not applicable
Inclusion criteria	Adult patients (aged >18 years) with symptomatic AS referred to hospital for consideration of valve surgery.
Exclusion criteria	Past myocardial infarction, more than mild mitral valve disease, previous cardiac surgery, patients in urgent need for surgery due to severe symptoms or heart failure, hypotension (systolic <110mmHg), current use of ACE inhibitors or ARBs, complicated diabetes, primary cardiomyopathy, potential pregnancy or breast-feeding, recent history of malignancy, history of alcohol or drug abuse, elevated serum creatinine (>176 micromol/L), and participation in another investigational drug study.
Recruitment/selection of patients	May 2009 - August 2012; consecutive patients screened
Age, gender and ethnicity	Age - Mean (SD): Intervention: 73±9, control: 70±12. Gender (M:F): 22:27. Ethnicity: Not stated
Further population details	1. Age: Mixed (Candesartan mean: 73+/-9, Placebo mean: 70+/-12). 2. Heart rate: Normal (Candesartan mean: 68+/-15, Placebo mean: 67+/-13). 3. Presence vs. absence of uncontrolled systemic hypertension: Mixed (Candesartan mean: 134/78+/-14/12 (could be classified as hypertensive towards the higher values). Placebo mean: 137/80+/-21/13). 4. Severe vs non-severe HVD: Severe 5. Symptomatic vs asymptomatic: Symptomatic
Extra comments	Severity: aortic valve area index (cm²/m²) - 0.42 (0.13) and 0.41 (0.11); mean LV-AO pressure gradient (mmHg): 52 (14) and 49 (14). Patients who did not meet the exclusion criteria and could be put on the hospital's normal waiting list for invasive investigations and surgery were included
Indirectness of population	No indirectness: Possible that the NYHA classes could not be completely applicable (possible selective reporting).
Interventions	(n=25) Intervention 1: Angiotensin-II receptor blockers (ARBs) - Candesartan. 8mg per day for 2 weeks, and then 16mg per day until 3 days before they have valve surgery. Duration Mean = 5.4 months, with some

	higher (up to 1 year) and some lower (2 months). Concurrent medication/care: Not reported. Indirectness: No indirectness  (n=26) Intervention 2: Placebo. Placebo. Duration Mean = 5.4 months, with some higher (up to 1 year) and some lower (2 months). Concurrent medication/care: Not reported. Indirectness: No indirectness
Funding	Equipment / drugs provided by industry (Also academic support (Finnish Foundation for Cardiovascular Research, EVO research funds of the Helsinki University Central Hospital, the Jenny and Antti Wihuri Foundation)

#### RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: CANDESARTAN versus PLACEBO

Protocol outcome 1: Hospitalisation due to heart failure at 12 months

- Actual outcome for Primary aortic [including bicuspid] stenosis: Acute heart failure and anaemia at Mean: 5.4 months. Range: 2-12 months; Group 1: 1/25, Group 2: 0/26; Comments: Defined as acute decompensated heart failure, so hospitalisation is assumed

Risk of bias: All domain - Very high, Selection - High, Blinding - Low, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: Serious indirectness, Comments: Possible indirectness dependent on whether the anaemia caused the heart failure; Baseline details: More people taking statins in the candesartan group. Higher proportion NYHA class III in placebo group (27% vs 12% in candesartan group).; Group 1 Number missing: 3, Reason: 2 due to adverse events, 1 due to being denied valve surgery; Group 2 Number missing: 5, Reason: 2 due to adverse events, 2 significant coronary artery disease, 1 preoperative exitus

Protocol outcome 2: Exercise tolerance at 12 months

- Actual outcome for Primary aortic [including bicuspid] stenosis: 6 minute walk test at Mean: 5.4 months. Range: 2-12 months; Group 1: mean -20 meters (SD 42); n=22, Group 2: mean -2 meters (SD 59); n=21; Comments: Baseline values:

Candesartan: 390 (99), Placebo: 380 (197). Final values are not reported separately.

Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: More people taking statins in the candesartan group. Higher proportion NYHA class III in placebo group (27% vs 12% in candesartan group). Group 1 Number missing: 3, Reason: 2 due to adverse events, 1 due to being denied valve surgery; Group 2 Number missing: 5, Reason: 2 due to adverse events, 2 due to significant coronary artery disease, 1 had preoperative exitus

Protocol outcome 3: Withdrawal due to adverse events at 6 months

- Actual outcome for Primary aortic [including bicuspid] stenosis: Withdrawal due to adverse events at Mean: 5.4 months. Range: 2-12 months; Group 1: 2/25, Group 2: 2/26; Comments: All due to dizziness

Risk of bias: All domain - Very high, Selection - High, Blinding - Low, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: More people taking statins in the candesartan group. It doesn't trigger their significant P value, but the NYHA class looks like the placebo group has more severe cases. Group 1 Number missing: 1, Reason: Denied valve surgery; Group 2 Number missing: 3, Reason: Significant coronary artery disease (2); preoperative exits (1)

- Actual outcome for Primary aortic [including bicuspid] stenosis : All-cause mortality at Mean: 5.4 months. Range: 2-12 months; Group 1: 0/25, Group 2: 1/26; Comments: Sudden death while awaiting surgery

Risk of bias: All domain - Very high, Selection - High, Blinding - Low, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: More people taking statins in the candesartan group. Higher proportion NYHA class III in placebo group (27% vs 12% in candesartan group). Group 1 Number missing: 3, Reason: 2 due to adverse events, 1 due to being denied valve surgery; Group 2 Number missing: 5, Reason: 2 due to adverse events, 2 due to significant coronary artery disease, 1 had preoperative exitus

Protocol outcomes not reported by the
study

Quality of life at 6 months; Cardiac mortality at 12 months; Quality of life at 12 months; Need for valve intervention at 6 months; Need for valve intervention at 12 months; Withdrawal due to adverse events at 12 months; Cardiac mortality at 6 months

Study	Klein 1985 <sup>87</sup>
Study type	RCT (Patient randomised; Crossover: Not reported)
Number of studies (number of participants)	1 (n=13)
Countries and setting	Conducted in South Africa; Setting: Unclear
Line of therapy	1st line
Duration of study	Intervention time: Two phases consisting of 2 weeks each (atenolol and placebo phases)
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: Diagnosis of mitral stenosis made clinically and confirmed by echocardiography. Mitral stenosis was considered 'significant' in every patient based on an echocardiographic finding of a mitral valve orifice area <1.5 cm.
Stratum	Primary mitral stenosis: All patients with mitral stenosis
Subgroup analysis within study	Not applicable
Inclusion criteria	Significant isolated mitral stenosis and sinus rhythm
Exclusion criteria	Not reported
Recruitment/selection of patients	Consecutive patients
Age, gender and ethnicity	Age - Range: 15-35 years. Gender (M:F): 2:11. Ethnicity: Not reported
Further population details	1. Age: <75 years (Age range of those included was 15-35 years). 2. Heart rate: Not stated / Unclear (States post-treatment values only). 3. Presence vs. absence of uncontrolled systemic hypertension: Not stated / Unclear (States post-treatment values only). 4. Severe vs non-severe HVD: Not stated / Unclear (Significant

	mitral stenosis defined as mitral valve orifice area <1.5 cm). 5. Symptomatic vs asymptomatic: Symptomatic (NYHA class II or III - corresponds to mild or moderate symptoms of heart failure).
Extra comments	Patients with isolated mitral stenosis and in sinus rhythm. Functional class II or III of the New York Heart Association classification. All had evidence of pulmonary arterial hypertension (based on palpable right ventricular impulse and a loud pulmonic component of the second heart sound).
Indirectness of population	Serious indirectness: Includes some patients under 18 years of age but the proportion is unclear. Mean age not reported.
Interventions	(n=13) Intervention 1: Beta blockers - Atenolol. 100 mg oral atenolol taken in the morning for 2 weeks. Duration 2 weeks. Concurrent medication/care: All patients received oral diuretic therapy throughout the study period. Indirectness: Serious indirectness; Indirectness comment: Unclear whether the treatment was first line  Comments: 6 patients had atenolol during first phase of crossover study and 7 patients had atenolol during second phase of crossover study  (n=13) Intervention 2: Placebo. Placebo taken in the morning for 2 weeks. Placebo tablet was identical in appearance to the atenolol tablet. Duration 2 weeks. Concurrent medication/care: All patients received oral diuretic therapy throughout the study period. Indirectness: Serious indirectness; Indirectness comment: Unclear whether treatment was first line  Comments: 7 patients had placebo during first phase of crossover study and 6 patients had placebo during second phase of crossover study
Funding	Equipment / drugs provided by industry (Atenolol and placebo tablets were provided by Imperial Chemical Industries, Inc.)

# RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: ATENOLOL versus PLACEBO

Protocol outcome 1: Exercise tolerance at 12 months

- Actual outcome for Primary mitral stenosis: Total duration of exercise (during modified Bruce protocol) at End of 2-week treatment period; Group 1: mean 11 minutes (SD 2); n=13, Group 2: mean 9 minutes (SD 2); n=13; Comments: Exercise testing was performed on commercial treadmills according to a modified Bruce protocol until the point of exhaustion, dizziness or severe dyspnoea was reached
- Risk of bias: All domain Very high, Selection Low, Blinding High, Incomplete outcome data Low, Outcome reporting Low, Measurement Low, Crossover High, Subgroups Low, Other 1 Low; Indirectness of outcome: Serious indirectness, Comments: Follow-up is end of study rather than 12 month follow-up; Baseline details: Crossover study same participants in both groups; Blinding details: Statement that atenolol and placebo tablets were identical in appearance; Group 1 Number missing: 0; Group 2 Number missing: 0
- Actual outcome for Primary mitral stenosis: Maximal exercise capacity (during modified Bruce protocol) at End of 2-week treatment period; Group 1: mean 84 work units (SD 47); n=13, Group 2: mean 45 work units (SD 29); n=13; Comments: Exercise testing was performed on commercial treadmills according to a modified Bruce protocol until the point of exhaustion, dizziness or severe dyspnoea was reached. Maximal exercise capacity derived from following equation, which was calculated for each stage of exercise and then summated: [time (min) x speed (km/h) x incline (degrees)/3 minutes]. The index takes into account the time spent exercising as well as the increasing difficulty of exercise with each successive stage.

Protocol outcomes not reported by the study

Quality of life at 6 months; Hospitalisation due to heart failure at 12 months; Cardiac mortality at 12 months; Quality of life at 12 months; Need for valve intervention at 6 months; Need for valve intervention at 12 months; Withdrawal due to adverse events at 6 months; Withdrawal due to adverse events at 12 months; All-cause mortality at 12 months; Cardiac mortality at 6 months

Study	Kumar 1994 <sup>90</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=31)
Countries and setting	Conducted in India; Setting: Secondary care
Line of therapy	1st line
Duration of study	Intervention + follow up: 6 months
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: Previously seen in clinic. Have an echo during the procedure to verify it. Baseline NYHA class recorded.
Stratum	Primary mitral stenosis
Subgroup analysis within study	Not applicable
Inclusion criteria	Isolated symptomatic rheumatic mitral stenosis in sinus rhythm
Exclusion criteria	Mitral valve area <0.8cm²; haemoglobin <12gm%; obstructive lung disease; >grade I mitral or aortic regurgitation on 2D echocardiogram and colour doppler
Recruitment/selection of patients	Consecutive
Age, gender and ethnicity	Age - Mean (SD): Intervention: 23.6±7.7; Placebo: 22.8±8.2. Gender (M:F): 15:11. Ethnicity: Not stated
Further population details	1. Age: <75 years (Metoprolol mean: 23.6+/-7.7; Placebo mean: 22.8+/-8.2). 2. Heart rate: Mixed (Metoprolol mean: 88.3+/-16.7, Placebo mean: 91.8+/-10.8. At the extreme high would be tachycardic). 3. Presence vs. absence of uncontrolled systemic hypertension: Absence of uncontrolled systemic hypertension (states that no one was recruited who had a systemic blood pressure >140/90mmHg). 4. Severe vs non-severe HVD: Not stated / Unclear 5. Symptomatic vs asymptomatic : Symptomatic
Extra comments	All had been advised surgery or balloon valvotomy and were awaiting intervention or had declined. The exclusion criteria are not very well defined (not prespecified, just noted in response to the patients they had)
Indirectness of population	No indirectness
Interventions	(n=13) Intervention 1: Beta blockers - Metoprolol. 25mg BD increasing up to 50mg BD dependent on patient preference. Duration 6 months. Concurrent medication/care: Not stated - it does state that digoxin was not given. Indirectness: No indirectness
	(n=13) Intervention 2: Placebo. Placebo. Duration 6 months. Concurrent medication/care: As previously - not stated. However, no patients were given digoxin. Indirectness: No indirectness
Funding	Funding not stated

Protocol outcome 1: Exercise tolerance at 12 months

- Actual outcome for Primary mitral stenosis: Pulmonary capillary wedge pressure after exercise

at 6 months; Group 1: mean 35.7 mmHg (SD 7.3); n=13, Group 2: mean 50.5 mmHg (SD 10.4); n=13; Comments: Baseline values: metoprolol - 40.3±10.8; placebo 34.1±10.6

Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: Serious indirectness, Comments: Surrogate outcome measure; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the study

Hospitalisation due to heart failure at 12 months; Cardiac mortality at 12 months; Quality of life at 12 months; Need for valve intervention at 6 months; Need for valve intervention at 12 months; Withdrawal due to adverse events at 6 months; Withdrawal due to adverse events at 12 months; All-cause mortality at 12 months; Cardiac mortality at 6 months

4 MIDs used to assess imprecision were ±4.0

disease: DRAFT FOR CONSULTATION

Study	Patel 1995 <sup>123</sup>								
Study type	RCT (Patient randomised; Crossover: 1 week)								
Number of studies (number of participants)	1 (n=19)								
Countries and setting	Conducted in South Africa; Setting: Secondary care								
Line of therapy	1st line								
Duration of study	Intervention + follow up: 4 weeks								
Method of assessment of guideline condition	Unclear method of assessment/diagnosis: Not stated.								
Stratum	Primary mitral stenosis:								
Subgroup analysis within study	Not applicable								
Inclusion criteria	People with symptomatic isolated tight pliable mitral stenosis admitted for percutaneous mitral valvotomy with NYHA class II-III heart failure, four of whom had a previous surgical closed mitral valvotomy.								
Exclusion criteria	Presence of atrial fibrillation, right heart failure, obstructive or embolic lung disease, and any contraindication to beta-blocker therapy.								
Recruitment/selection of patients	From their patients admitted to hospital								
Age, gender and ethnicity	Age - Mean (range): 28 (17-51). Gender (M:F): 3:16. Ethnicity: Not stated								
Further population details	1. Age: <75 years (Range = 17-51). 2. Heart rate: Normal (82+/-11). 3. Presence vs. absence of uncontrolled systemic hypertension: Not stated / Unclear 4. Severe vs non-severe HVD: Not stated / Unclear 5. Symptomatic vs asymptomatic: Symptomatic								
Extra comments	All medication except maintenance diuretic therapy was discontinued for at least 7 days prior to enrolment.								
Indirectness of population	No indirectness								
Interventions	(n=19) Intervention 1: Beta blockers - Acebutolol. Acebutolol 400mg daily or Atenolol 100mg daily. Duration 1 week. Concurrent medication/care: Background diuretic therapy, but all others were stopped at least 7 days before the study protocol started. Indirectness: Serious indirectness; Indirectness comment: See atenolol - people in the intervention were given either acebutalol or atenolol with no way to tell which patient received which in the reported data.								
	(n=19) Intervention 2: Placebo. Placebo. Duration 1 week. Concurrent medication/care: Background diuretic therapy, but all others were stopped at least 7 days before the study protocol started. Indirectness: No indirectness								
Funding	Funding not stated								

## RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: ACEBUTOLOL/ATENOLOL versus PLACEBO

Protocol outcome 1: Exercise tolerance at 12 months

- Actual outcome for Primary mitral stenosis: Treadmill exercise time to exhaustion (using Weber's protocol) at 4 weeks; Group 1: mean 8.8 min (SD 1.7); n=19, Group 2: mean 9.4 min (SD 1.8); n=19; Comments: Baseline value = 9.2±1.8

Risk of bias: All domain - High, Selection - Very high, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: Reports baseline only for the parameters they are measuring; Group 1 Number missing: ; Group 2 Number missing:

- Actual outcome for Primary mitral stenosis: Heart rate at peak exercise at 4 weeks; Group 1: mean 63 beats per minute (SD 10); n=19, Group 2: mean 78 beats per minute (SD 9); n=19

Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: Serious indirectness, Comments: Time period is only 4 weeks rather than 12 months; Baseline details: Reports baseline only for the parameters they are measuring but crossover trial so will be matched; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the study

Quality of life at 6 months; Hospitalisation due to heart failure at 12 months; Cardiac mortality at 12 months; Quality of life at 12 months; Need for valve intervention at 6 months; Need for valve intervention at 12 months; Withdrawal due to adverse events at 6 months; Withdrawal due to adverse events at 12 months; All-cause mortality at 12 months; Cardiac mortality at 6 months

4 MIDs used to assess imprecision were ±4.0

disease: DRAFT

FOR

CONSULTATION

Study	Seneviratne 1994 <sup>147</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=28)
Countries and setting	Conducted in Australia; Setting: Secondary care
Line of therapy	1st line
Duration of study	Intervention + follow up: 12 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: Echocardiography and clinical assessment
Stratum	Secondary heart valve disease – mitral regurgitation or tricuspid regurgitation: Mitral regurgitation secondary to dilated heart failure
Subgroup analysis within study	Not applicable
Inclusion criteria	NYHA grade II-III with dilated left ventricles, ejection fraction <40% and a functional mitral regurgitant area of >5cm².
Exclusion criteria	Myocardial infarction within the preceding three months; unstable or severe angina pectoris; valvar (?valvular) heart disease; serum creatinine >0.18mmol/L, a history of alcohol misuse, and ACE inhibitor treatment of either >25mg of captopril or >5mg of enalapril a day.
Age, gender and ethnicity	Age - Mean (range): 71.6 (57-80). Gender (M:F): 27:1. Ethnicity: Not stated
Further population details	1. Age: Mixed (Captopril mean = 72.3+/-5.4, Placebo mean = 71.5+/-7.2 - Fall under 75 for the mean, but confidence intervals cross. Therefore, mixed?). 2. Heart rate: Not stated / Unclear 3. Presence vs. absence of uncontrolled systemic hypertension: Absence of uncontrolled systemic hypertension (arterial pressure reported (rather than systolic/diastolic). Captopril mean = 99+/-13.8, Placebo mean = 93+/-9.8). 4. Severe vs non-severe HVD: Not stated / Unclear (No severity mentioned). 5. Symptomatic vs asymptomatic: Symptomatic (NYHA class II or III).
Extra comments	Severity of regurgitation not stated
Indirectness of population	No indirectness
Interventions	(n=14) Intervention 1: Angiotensin-converting enzyme (ACE) inhibitors - Captopril. 6.25mg twice daily, increasing to 12.5mg twice daily after 4 weeks, increasing to 25mg twice daily after 8 weeks, increasing to 50mg twice daily at 12 weeks. Duration 6 months. Concurrent medication/care: Randomisation occurred after a 2 week placebo washout period and a test dose of 6.25 mg oral captopril. Digoxin, diuretics and nitrates were continued. Indirectness: No indirectness
	(n=14) Intervention 2: Placebo. Placebo. Duration 6 months. Concurrent medication/care: Randomisation occurred after a 2 week placebo washout period and a test dose of 6.25 mg oral captopril. Digoxin, diuretics

Funding Study funded by industry (Aid of a grant from Bristol-Squibb-Myers who played no part in the analysis and interpretation of the data.)		and nitrates were continued. Indirectness: No indirectness
	Funding	

## RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: CAPTOPRIL versus PLACEBO

Protocol outcome 1: Quality of life at 6 months

- Actual outcome for Secondary heart valve disease – mitral regurgitation or tricuspid regurgitation: Duke activity index score at 12 weeks; Group 1: mean 29 No units (SD 8.9); n=10, Group 2: mean 22.3 No units (SD 9.8); n=13; Comments: Baseline characteristics:

Captopril Duke activity status index: 21.5 (7.8)

Placebo Duke activity status index: 22.6 (11)

Risk of bias: All domain - Very high, Selection - High, Blinding - Low, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: Serious indirectness, Comments: Only measures physical activity rather than other aspects of quality of life; Group 1 Number missing: 4; Group 2 Number missing: 1

Protocol outcome 2: Cardiac mortality at 6 months

- Actual outcome for Secondary heart valve disease – mitral regurgitation or tricuspid regurgitation: Cardiac mortality at 12 weeks; Group 1: 0/14, Group 2: 1/14; Comments: Acute MI

Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 0; Group 2 Number missing: 0

Protocol outcome 3: Withdrawal due to adverse events at 6 months

- Actual outcome for Secondary heart valve disease – mitral regurgitation or tricuspid regurgitation: Withdrawals due to adverse events at 12 weeks; Group 1: 0/14, Group 2: 0/13

Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 0; Group 2 Number missing: 0

Protocol outcomes not reported by the study

Cardiac mortality at 12 months; Quality of life at 12 months; Exercise tolerance at 12 months; Need for valve intervention at 6 months; Need for valve intervention at 12 months; Withdrawal due to adverse events at 12 months; All-cause mortality at 12 months; Hospitalisation due to heart failure at 12 months

Study	Shu 2005 <sup>150</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=88)
Countries and setting	Conducted in China; Setting: Secondary care
Line of therapy	1st line
Duration of study	Intervention + follow up: 12 months
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: Clinical assessment and echocardiographic evidence
Stratum	Primary mitral stenosis: Significant mitral stenosis with or without accompanying mitral valve regurgitation or aortic lesions (as a result of rheumatic heart disease)
Subgroup analysis within study	Not applicable
Inclusion criteria	1) History of uncorrected rheumatic heart valvular disease or NYHA class III or IV disease necessitating hospitalisation; 2) cardiothoracic ratio of less than 65%; 3) AF with a resting ventricular rate of 70 beats/minute or more for at least 3 months, as depicted by ECG; 4) An echocardiogram showing a significant mitral stenosis or aortic lesions and mitral valve regurgitation
Exclusion criteria	Uncorrected congenital heart disease; sustained ventricular tachycardia; severe liver and kidney dysfunction; chronic obstructive pulmonary disease; bronchial asthma; obstructive or restrictive cardiomyopathy or myocarditis; myocardial infarction; or unstable angina within the past three months. Additionally if they required intensive care or concurrent IV therapy; or were using calcium channel blockers, class I or III antiarrhythmic drugs, MAO-inhibitors or beta2-agonists.
Recruitment/selection of patients	Recruitment from their patients. No obvious consecutive patients.
Age, gender and ethnicity	Age - Mean (SD): Intervention = 40.6±6.8; Control = 43.5±7.4. Gender (M:F): 24:43. Ethnicity: Not stated
Further population details	1. Age: <75 years (Control mean = 43.5+/-7.4, Treatment mean = 40.6+/-6.8). 2. Heart rate: Mixed (Gives ventricular rate (which would indicate mixed between normal and tachycardic). Is this equivalent to heart rate? Control mean = 105+/-19, Treated mean = 110+/-21). 3. Presence vs. absence of uncontrolled systemic hypertension: Absence of uncontrolled systemic hypertension (systolic BP given. Control mean = 121+/-14, Treated mean = 115+/-12). 4. Severe vs non-severe HVD: Not stated / Unclear (Severity not mentioned). 5. Symptomatic vs asymptomatic: Symptomatic (NYHA class III or IV during admission to hospital).
Indirectness of population	No indirectness: Patients have multiple types of valve lesion at the same time. All patients had atrial fibrillation.
Interventions	(n=44) Intervention 1: Beta blockers - Bisoprolol. Initial dose of 1.25mg/day. Recommended maximum dose of 10mg/day. Gradual titration over 3 to 5 days by 2 to 3 weeks. Duration 6 to 12 months. Concurrent

	medication/care: All patients received basic therapy using one of the following: a diuretic, digoxin (extracted from Digitalis lanata), ACE-inhibitors (or ARBs if ACE-inhibitors were not tolerated) or nitrates. All received warfarin. Indirectness: No indirectness  (n=44) Intervention 2: Usual care. All patients received basic therapy using one of the following: a diuretic, digoxin (extracted from Digitalis lanata), ACE-inhibitors (or ARBs if ACE-inhibitors were not tolerated) or nitrates. All received warfarin. Duration 6 to 12 months. Concurrent medication/care: N/A. Indirectness: No
	indirectness
Funding	Academic or government funding (One of the doctors was supported by a fellowship from the Departments of Cardiology and Ultrasound diagnosis in Southwest Hospital)

### RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: BISOPROLOL versus USUAL CARE

Protocol outcome 1: Hospitalisation due to heart failure at 12 months

- Actual outcome: Hospitalisation due to exacerbated heart failure at 12 months; Group 1: 3/33, Group 2: 10/34

Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Blinding details: Even though the measures are objective, there is still a large risk of care being different due to the intervention not being blinded; Group 1 Number missing: 11, Reason: Adverse drug reactions in 5. Then they provide overall information - 14 excluded from evaluation at follow up with 7 having insufficient quality ECG/echo data, and 7 having poor telephone-connection difficulties. From the bits reported, the numbers don't add up so there is not full reporting; Group 2 Number missing: 10, Reason: See experimental group.

Protocol outcome 2: Exercise tolerance at 12 months

- Actual outcome: 6-minute walking distance at 6-12 months; Group 1: mean 423 meters (SD 25); n=33, Group 2: mean 290 meters (SD 23); n=34; Comments: Values at hospital discharge:

Treated =  $391\pm32$ 

Control =  $309\pm28$ 

Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low, Crossover - Low, Comments - ; Indirectness of outcome: No indirectness ; Blinding details: Even though the measures are objective, there is still a large risk of care being different due to the intervention not being blinded; Group 1 Number missing: 11, Reason: Adverse drug reactions in 5. Then they provide overall information - 14 excluded from evaluation at follow up with 7 having insufficient quality ECG/echo data, and 7 having poor telephone-connection difficulties. From the bits reported, the numbers don't add up so there is not full reporting; Group 2 Number missing: 10, Reason: See experimental group.

Protocol outcome 3: Withdrawal due to adverse events at 12 months

- Actual outcome: Withdrawal due to adverse events at 12 months; Group 1: 5/44, Group 2: 0/44; Comments: They don't report all of the withdrawal reasons for each patient (the figures don't add up) - a little dubious. Also don't report if any withdrawals due to drug effects in control group. Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - High, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Blinding details: Even though the measures are objective, there is still a large risk of care being different due to the intervention not being blinded; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the
study

Quality of life at 6 months; Cardiac mortality at 12 months; Quality of life at 12 months; Need for valve intervention at 6 months; Need for valve intervention at 12 months; Withdrawal due to adverse events at 6 months; All-cause mortality at 12 months; Cardiac mortality at 6 months

# **Appendix E: Forest plots**

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# E.4 Valve disease without heart failure

# E.14 Primary aortic [including bicuspid] stenosis

# E.1.151 ACE inhibitors compared to placebo

# Figure 3: Need for heart valve intervention at 12 months

	ACE-inhibitors Placebo Risk Ratio						Risk Ratio					
Study or Subgroup	Events	Total	Events	Total	M-H, Fixed, 95% CI			M-H, Fix	ed, 95%	CI		
Bull 2015	4	40	2	43	2.15 [0.42, 11.10]				<del>                                     </del>		<u> </u>	
						0.1	0.2	0.5	$\frac{1}{1}$	5	10	
							Fa	vours ACE-I	Favour	s placebo		

6

Figure 4: Exercise tolerance (exercise distance measured with treadmill exercise test, meters, change score) at 12 months

	ACE-inhibitors			ACE-inhibitors Placebo Mean Difference						Mean Difference						
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	IV, Fixed, 95% CI		IV, Fixe	d, 95% CI						
Bull 2015	-20	26	26	29	25	41	-49.00 [-61.59, -36.41]		<del>-</del>							
								-100	-50	0 5						

MIDs used to assess imprecision were calculated by multiplying the median baseline SD across study arms (373) by 0.5 and were ±187.0.

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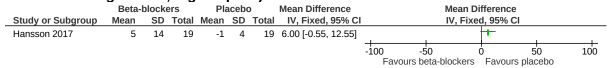
Figure 5: Withdrawal due to adverse events at 12 months

J	ACE-inhi	ors Placebo Risk Ratio					Risk Ratio					
Study or Subgroup	Events	Total	<b>Events</b>	Total	M-H, Fixed, 95% CI							
Bull 2015	2	38	1	42	2.21 [0.21, 23.41]				<del>                                     </del>			
						0.01	0.	1	1	10	100	
							Fav	ours ACE-I	Favours p	lacebo		

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#### E.1.1112 Beta blockers compared to placebo

Figure 6: Quality of life (Minnesota living with heart failure questionnaire, 0-105, change score, high is poor) at 5 months



12 Published MIDs of ±5.0 were used to assess imprecision for MLWHF questionnaire.

13

Figure 7: Exercise tolerance (6 minute walk test distance, meters, change score) at 5 months

	Beta-l	block	ers	Pla	aceb	0	Mean Difference		M	ce		
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	IV, Fixed, 95% CI		I۱	/, Fixed, 95%	CI	
Hansson 2017	2	46	19	14	49	19	-12.00 [-42.22, 18.22]			-		
								-100	-50	Ó	50	100
									Favours pla	acebo Favo	irs beta-block	ers

MIDs used to assess imprecision were calculated by multiplying the median baseline SD across study arms (41) by 0.5 and were ±21.0.

Figure 8: Withdrawal or dose reduction due to adverse events at 5 months

	Beta-bloc	ckers	Placel	00	RISK Ratio			RISK	Ratio			
Study or Subgroup	Events	Total	<b>Events</b>	Total	M-H, Fixed, 95% CI			M-H, Fixe	ed, 95%	CI		
Hansson 2017	4	19	2	19	2.00 [0.41, 9.65]							
						0.1 0.2 0.5 1 2 5					5	10
						F	avours be	eta-blockers	Favour	s placebo		

# E.1.153 Diuretics compared to placebo

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Figure 9: All-cause mortality at 19 months

	Diuretics Placebo Risk Ratio						Risk Ratio							
Study or Subgroup	Events	Total	<b>Events</b>	Total	M-H, Fixed, 95% CI			M-H, Fiz	xed,					
Stewart 2008	1	30	2	31	0.52 [0.05, 5.40]	<del> </del>		+	$\top$					
						0.1	0.2	0.5	1_	2 avours place	5	10		

Figure 10: Cardiac mortality at 19 months

_	Diuretics		Placebo		Peto Odds Ratio	Peto Odds Ratio					
Study or Subgroup	Events	Total	Events	Total	Peto, Fixed, 95% CI		Peto, Fixed, 95% CI				
Stewart 2008	0	29	1	30	0.14 [0.00, 7.06]	+		1			
						0.01	0.	1	1 10	100	
							Favou	rs diuretics	Favours place	ebo	

Figure 11: Quality of life (SF-36 physical functioning subscale, 0-100, change score, high is good) at 12 months

	Diuretics			Placebo			Mean Difference		Mean Difference			
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	IV, Fixed, 95% CI		IV	Fixed, 95%	CI	
Stewart 2008	<b>-</b> 5	22	29	-9	19	30	4.00 [-6.50, 14.50]			+		
								-100	-50	Ó	50	100
									Favours placebo Favours diuretics			

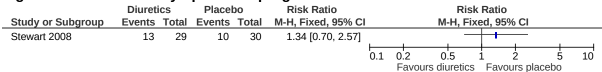
8 Published MIDs of ±3.0 were used to assess imprecision for SF-36 physical functioning subscale.

Figure 12: Quality of life (SF-36 role physical subscale, 0-100, change score, high is good) at 12 months

_	Diuretics			Placebo			Mean Difference	Mean	Mean Difference			
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	IV, Fixed, 95% CI		IV, Fix	ed, 95% CI		
Stewart 2008	-9	34	29	-12	37	30	3.00 [-15.12, 21.12]		_	+		
								-100	-50 Favours placebo	0 Favours o	50 diuretics	100

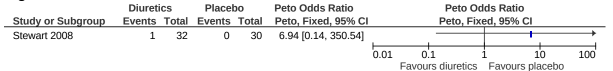
10 Published MIDs of ±3.0 were used to assess imprecision for SF-36 role physical subscale.

Figure 13: Onset of symptoms or progression of NYHA class at 19 months



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Figure 14: Withdrawal due to adverse events at 19 months



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### E.1.144 Statins compared to placebo

Figure 15: All-cause mortality at 4.3 years

19410 101 711 00	Statir		Placebo		,	Risk Ratio	Risk Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% C	CI M-H, Fixed, 95% CI	
Chan 2010 (ASTRONOMER)	3	134	5	135	4.7%	0.60 [0.15, 2.48]	_ <del>-</del>	
Dichtl 2008 (TASS)	1	24	1	23	1.0%	0.96 [0.06, 14.43]	<u> </u>	
Rossebo 2008 (SEAS)	105	944	100	929	94.4%	1.03 [0.80, 1.34]	i 📮	
Total (95% CI)		1102		1087	100.0%	1.01 [0.79, 1.30]	<b>•</b>	
Total events	109		106					
Heterogeneity: Chi <sup>2</sup> = 0.54, df =	2 (P = 0.7	'6); I <sup>2</sup> =	0%				0.01 0.1 1 10 1	100
Test for overall effect: $Z = 0.10$	(P = 0.92)						Favours statin Favours placebo	100

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Figure 16: All-cause mortality at 4.4 years (time-to-event data)

			Hazard Ratio			Haza	ıra Ka	tio		
Study or Subgroup	log[Hazard Ratio]	SE	IV, Fixed, 95% CI			IV, Fix	ed, 95	% CI		
Rossebo 2008 (SEAS)	0.0392	0.1403	1.04 [0.79, 1.37]			-	+			
			,	0.1	0.2	0.5	1	2	5	10
					Fa	vours statir	n Fav	ours p	lacebo	

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Figure 17: Cardiac mortality at 3.7 years

	Stati	n	Placebo			Risk Ratio		Risk Ratio			
Study or Subgroup	Events	Total	<b>Events</b>	Total	Weight	M-H, Fixed, 95% C	I	M-H, Fixe	ed, 95% CI		
Chan 2010 (ASTRONOMER)	2	134	5	135	6.1%	0.40 [0.08, 2.04]					
Cowell 2005 (SALTIRE)	11	77	19	78	23.2%	0.59 [0.30, 1.15]			+		
Dichtl 2008 (TASS)	1	24	1	23	1.3%	0.96 [0.06, 14.43]					
Rossebo 2008 (SEAS)	47	944	56	929	69.4%	0.83 [0.57, 1.20]		-	-		
Total (95% CI)		1179		1165	100.0%	0.75 [0.54, 1.03]		•			
Total events	61		81								
Heterogeneity: Chi <sup>2</sup> = 1.36, df =	3 (P = 0.7)	72); I <sup>2</sup> =	0%				-		<u> </u>	100	
Test for overall effect: Z = 1.80 (	P = 0.07	,,					0.01	0.1 Favours statin	1 10 Favours placebo	100	

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Figure 18: Cardiac mortality at 4.4 years (time-to-event data)

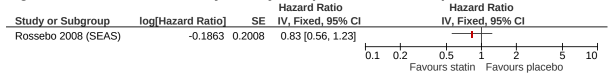


Figure 19: Onset of symptoms or progression of NYHA class at 3.2 years



Figure 20: Onset of symptoms of progression of NYHA class at 4.4 years (time-to-event data)

			Hazard Ratio			Hazard	d Ratio			
Study or Subgroup	log[Hazard Ratio]	SE	IV, Fixed, 95% CI			IV, Fixed	d, 95%	CI		
Rossebo 2008 (SEAS)	0.0862	0.2879	1.09 [0.62, 1.92]				1			
				0.1	0.2	0.5	1 :	<del> </del> 2	5	10
					Fa	vours statin	Favou	rs placeb	0	

Figure 21: Need for heart valve intervention at 3.7 years

_	Stati	n	Place	bo	Risk Ratio			Risk Ratio
Study or Subgroup	Events	Total	<b>Events</b>	Total	Weight	M-H, Random, 95% CI		M-H, Random, 95% CI
Chan 2010 (ASTRONOMER)	28	134	27	135	24.1%	1.04 [0.65, 1.67]		<del>-</del>
Cowell 2005 (SALTIRE)	11	77	19	78	14.3%	0.59 [0.30, 1.15]		<del></del>
Dichtl 2008 (TASS)	5	25	1	24	1.8%	4.80 [0.60, 38.14]		<del>                                     </del>
Rossebo 2008 (SEAS)	267	944	278	929	59.7%	0.95 [0.82, 1.09]		•
Total (95% CI)		1180		1166	100.0%	0.93 [0.70, 1.24]		•
Total events	311		325					
Heterogeneity: Tau <sup>2</sup> = 0.03; Chi <sup>2</sup>	$^2 = 4.47$ , d	lf = 3 (F	P = 0.22;	$I^2 = 33^\circ$	%		0.01	0.1 1 10 100
Test for overall effect: Z = 0.48 (	P = 0.63)						0.01	0.1 1 10 100 Favours statin Favours placebo

Figure 22: Need for heart valve intervention at 4.4 years (time-to-event data)

			Hazard Ratio			Haza	ra Ka	itio		
Study or Subgroup	log[Hazard Ratio]	SE	IV, Fixed, 95% CI			IV, Fix	ed, 9	5% CI		
Rossebo 2008 (SEAS)	0	0.089	1.00 [0.84, 1.19]		_		+			
				0.1	0.2	0.5	1	2	5	10
				0	Fav	ours statir	Fa	vours pla	acebo	

Figure 23: Withdrawal due to adverse events at 6 months

	Statin		Placebo		Peto Odds Ratio	Peto Odds Rat	io
Study or Subgroup	Events Total Events Total		Peto, Fixed, 95% CI	Peto, Fixed, 95%	6 CI		
Dichtl 2008 (TASS)	1	25	0	23	6.82 [0.13, 344.93]		<del></del>
						0.01 0.1 1 Favours statin Favou	10 100 rs placebo

Figure 24: Withdrawal due to adverse events at 3.3 years

_	Stati	n	Place	bo		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	<b>Events</b>	Total	Weight	M-H, Fixed, 95% C	CI M-H, Fixed, 95% CI
Chan 2010 (ASTRONOMER)	25	134	26	135	17.0%	0.97 [0.59, 1.59]	] —
Cowell 2005 (SALTIRE)	7	77	4	78	2.6%	1.77 [0.54, 5.81]	] -
Rossebo 2008 (SEAS)	144	943	122	929	80.4%	1.16 [0.93, 1.45]	i <del>-</del>
Total (95% CI)		1154		1142	100.0%	1.15 [0.94, 1.40]	1
Total events	176		152				
Heterogeneity: Chi <sup>2</sup> = 0.98, df =	2 (P = 0.6	51); I <sup>2</sup> =	0%				
Test for overall effect: $Z = 1.33$	(P = 0.18)						0.1 0.2 0.5 1 2 5 10  Favours statin Favours placebo

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# E.12 Primary aortic regurgitation

### E.1.241 ACE inhibitors compared to placebo/no treatment

Figure 25: All-cause mortality at 7 years

	ACE-inhibitors		ACE-inhibitors No treatment							
Study or Subgroup	Events Total		<b>Events</b>	Total	M-H, Fixed, 95% CI		M-I	H, Fixed, 95	% CI	
Evangelista 2005	1	32	1	31	0.97 [0.06, 14.82]					
						0.01	0.1	1	10	100
							Favours /	ACE-I Favo	urs no treati	ment

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Figure 26: Cardiac mortality at 7 years

	ACE-inni	ACE-inhibitors		ment	Peto Odds Ratio		Pe	itio			
Study or Subgroup	Events Total		Events	Total	Peto, Fixed, 95% CI		Peto	, Fixed, 95	ed, 95% CI		
Evangelista 2005	0	32	1	31	0.13 [0.00, 6.61]	<del></del>					
					C	0.01	0.1	1	10	100	
							Favours A	CF-L Favo	ours no treatr	nent	

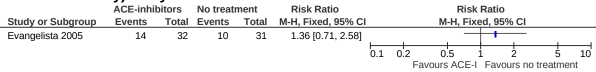
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Figure 27: Onset of symptoms or progression of NYHA class at 0.5-7 years

_	ACE-inhibitors			ACE-inhibitors No treatment			ment		Risk Difference	Risk Difference
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI			
Evangelista 2005	10	32	8	31	76.1%	0.05 [-0.17, 0.28]	<del></del>			
Wisenbaugh 1994	0	11	0	9	23.9%	0.00 [-0.18, 0.18]	<del>-</del>			
Total (95% CI)		43		40	100.0%	0.04 [-0.13, 0.22]				
Total events	10		8							
Heterogeneity: Chi2 =	0.23, df = 1	(P = 0.6)	3); $I^2 = 0\%$			<u>├</u> -1	-0.5 0 0.5 1			
Test for overall effect: Z = 0.46 (P = 0.64)						=1	Favours ACF-I Favours no treatment			

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Figure 28: Evidence of HVD progression on imaging (worsening of disease severity) at 7 years





	ACE-inhi	CE-inhibitors No treatment			Risk Ratio Risl				sk Ratio			
Study or Subgroup	Events	Total	<b>Events</b>	Total	M-H, Fixed, 95% CI			M-H, Fixe	ed, 95%	CI		
Evangelista 2005	16	32	12	31	1.29 [0.74, 2.27]				-	_		
						0.1	0.2	0.5	1 2	2	5	10
								avoure ACE-I	Favou	re no tra	atmai	nt .

Figure 30: Withdrawal due to adverse events at 7 years

	ACE-inhi	bitors	No treat	ment	Peto Odds Ratio		Peto	Odd	s Ratio		
Study or Subgroup	Events Total		<b>Events</b>	Total	l Peto, Fixed, 95% Cl		Peto,	Fixed	ed, 95% CI		
Evangelista 2005	3	32	0	31	7.65 [0.77, 76.34]		1	+			
						0.01	0.1	1	10	100	
							Favoure AC	`F_I	Favoure no treatr	mant	

2

# E.1.232 ACE inhibitors compared to calcium channel blockers

Figure 31: All-cause mortality at 7 years

_	ACE-inhi	bitors	Calcium-channel	blocker	Risk Ratio			Risk F	Ratio	
Study or Subgroup	Events	Total	Events	Total	M-H, Fixed, 95% CI		M-	H, Fixe	d, 95% CI	
Evangelista 2005	1	32	1	32	1.00 [0.07, 15.30]		<del></del>			_
						0.01	0.1	1	10	100
							For tours	ACE I	Covering coloium	abannal blaakar

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Figure 32: Cardiac mortality at 7 years

	ACE-inni	bitors	Calcium-channel	biocker	Peto Odds Ratio			Peto Od	ids Ratio		
Study or Subgroup	Events	Total	Events	Total	Peto, Fixed, 95% CI		F	Peto, Fixe	ed, 95% CI		
Evangelista 2005	0	32	1	32	0.14 [0.00, 6.82]	+					
						0.01	0.1		1	10	100
							Favou	rs ACF-I	Favours cale	cium-cha	nnel blocker

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Figure 33: Onset of symptoms or progression of NYHA class at 4.8 years

_	ACE-inhi	bitors	Calcium-channel blocker		Risk Difference		Risk Difference
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
Banaszewski 1998	0	12	0	13	28.1%	0.00 [-0.14, 0.14]	<del>-</del>
Evangelista 2005	10	32	8	32	71.9%	0.06 [-0.16, 0.28]	<del>-  </del>
Total (95% CI)		44		45	100.0%	0.04 [-0.12, 0.21]	
Total events	10		8				
Heterogeneity: Chi <sup>2</sup> = 0	0.41, df = 1	(P = 0.52)	2); I <sup>2</sup> = 0%			<u> </u>	-0.5 0 0.5 1
Test for overall effect:	Z = 0.54 (P	= 0.59)				-1	Favours ACE-I Favours calcium-channel blocker

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Figure 34: Evidence of HVD progression on imaging (worsening of disease severity) at 4.8 years

	, ,		,				
	ACE-inhi	bitors	Calcium-channel b	olocker		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	CI M-H, Fixed, 95% CI
Banaszewski 1998	0	13	2	12	20.6%	0.19 [0.01, 3.52]	1
Evangelista 2005	14	32	10	32	79.4%	1.40 [0.73, 2.67]	j <del> </del>
Total (95% CI)		45		44	100.0%	1.15 [0.62, 2.13]	•
Total events	14		12				
Heterogeneity: Chi <sup>2</sup> = '	1.83, df = 1	(P = 0.13)	8); I <sup>2</sup> = 45%				0.01 0.1 1 10 100
Test for overall effect:	7 _ 0 // /D	- 0.66)					
rest for overall effect.	Z = 0.44 (F	- 0.00)					Favours ACE-I Favours calcium-channel blocker



	ACE-inhi	bitors	Calcium-channel	blocker	Risk Ratio			Risk	k Ratio			
Study or Subgroup	Events	Total	Events	Total	M-H, Fixed, 95% CI			M-H, Fixe	ed, 95% C	l		
Evangelista 2005	16	32	13	32	1.23 [0.71, 2.12]		, <del>     </del>			-		
						0.1	0.2	0.5	1 2	2	5	10
								Egyourg ACE I	Egyoure	anlaium .	ohonnol	blookor

Figure 36: Withdrawal due to adverse events at 7 years

	ACE-inhi	bitors	Calcium-channel	blocker	Risk Ratio	Risk Ratio						
Study or Subgroup	Events	Total	Events	Total	M-H, Fixed, 95% CI			M-H, Fi	xed, 95%	CI		
Evangelista 2005	3	32	7	32	0.43 [0.12, 1.51]			<del></del>				
						0.1 0	.2	0.5	1	2	5	10
							F	avours ACF.	-I Favou	e calci	ıım-chann	al blocker

### E.1.243 ARBs compared to beta blockers

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# Figure 37: Exercise tolerance (exercise work rate using an ergometer, Watts, final value) at 3 weeks

	A	ARB		Beta-	block	er	Mean Difference		Me	ean Differen	ce	
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	IV, Fixed, 95% CI		IV	, Fixed, 95%	CI	
Roberts 2018	29	6	17	29	8	17	0.00 [-4.75, 4.75]	ı				
								-10	-5	0 ARB Favo	5	10

MIDs used to assess imprecision were calculated by multiplying the median control group final value SD across studies (8.0) by 0.5 and were ±4.0.

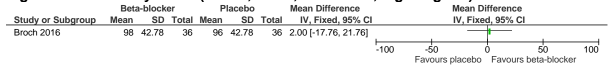
#### E.1.284 Beta blockers compared to placebo

# Figure 38: Quality of life (EuroQol visual analogue scale, 0-100, final value, high is good) at 6 months

	Beta-	-block	cer				Mean Difference		Me	ean Differen	ce	
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	IV, Fixed, 95% CI		IV	, Fixed, 95%	CI	
Broch 2016	85	7	36	82	16	36	3.00 [-2.70, 8.70]			+		
								-100	-50	Ó	50	100
									Favours pla	cebo Favo	urs beta-block	ker

9 MIDs used to assess imprecision were calculated by multiplying the median baseline SD across study arms (10.0) by 0.5 and were ±5.0.

Figure 39: Quality of life (KCCQ, 0-100, final value, high is good) at 6 months



MIDs used to assess imprecision were calculated by multiplying the median control group final value SD across studies (42.78) by 0.5 and were ±21.39.

### Figure 40: Exercise tolerance (peak work, watts, final value) at 6 months

	Beta-	-block	cer	Pla	acebo	0	Mean Difference		Mean I	Difference		
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	IV, Fixed, 95% CI		IV, Fix	ed, 95% CI		
Broch 2016	229	62	36	241	62	36	-12.00 [-40.64, 16.64]		<del></del>	<del>                                     </del>		
								-100	-50 Favours placebo	0 Favours m	50	100

MIDs used to assess imprecision were calculated by multiplying the median baseline SD across study arms (63.0) by 0.5 and were ±31.5.

### E.1.245 Calcium channel blockers compared to placebo/no treatment

Figure 41: All-cause mortality at 7 years

			No treat	ment	Risk Ratio		Risk Ratio			
Study or Subgroup	Events	Total	Events	Total	M-H, Fixed, 95% CI		M-H, Fix	ed, 95% CI		
Evangelista 2005	1	32	1	31	0.97 [0.06, 14.82]	_				
						0.01	.1	1 10	100	
						Favours calcium-	channel blocker	Favours no treatment		

Figure 42: Cardiac mortality at 7 years



Figure 43: Onset of symptoms or progression of NYHA class at 7 years

	Calcium-chamier b	IOCKCI	NO treat	IIIGIIL	Misk Matio				italio			
Study or Subgroup	Events	Total	Events	Total	M-H, Fixed, 95% CI			M-H, Fix	ed, 95% Cl			
Evangelista 2005	8	32	8	31	0.97 [0.42, 2.26]				•	_		
						0.1	0.2	0.5	1 :	2 :	5	10
						Favou	urs calcium	channel blocker	Favours r	no treatment		

Figure 44: Evidence of HVD progression on imaging (worsening of disease severity) at 7 years

	Calcium-channel blocker		No treat	ment	Risk Ratio	Risk Ratio							
Study or Subgroup	Events	Total	Events	Total	M-H, Fixed, 95% CI			M-H, Fix	ed, 95% C				
Evangelista 2005	10	32	10	31	0.97 [0.47, 2.00]				<u> </u>	-			
						0.1	0.2	0.5	1 :	2	5	10	
						Favou	urs calcium-	channel blocker	Favours i	no treatment			

Figure 45: Need for heart valve intervention at 1-7 years

							- <b>,</b>
_	Calcium-channel	blocker	No treat	ment		Risk Difference	Risk Difference
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% C	M-H, Fixed, 95% CI
Evangelista 2005	13	32	12	31	46.7%	0.02 [-0.22, 0.26]	<del></del>
Scognamiglio 1990	0	36	0	36	53.3%	0.00 [-0.05, 0.05]	+
Total (95% CI)		68		67	100.0%	0.01 [-0.11, 0.13]	•
Total events	13		12				
Heterogeneity: Chi2 =	0.12, df = 1 (P = 0.73	3); I <sup>2</sup> = 0%					-1 -0.5 0 0.5 1
Test for overall effect:	Z = 0.15 (P = 0.88)						Favours calcium-channel blocker Favours no treatment

9

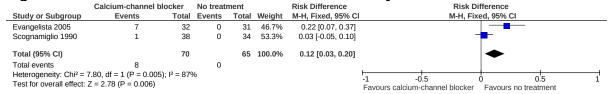
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## E.1.216 Digoxin compared to calcium channel blockers

Figure 47: All-cause mortality at 6 years

_	Digo	in	Calcium channel b	olocker	Peto Odds Ratio		Peto O	dds Ratio	
Study or Subgroup	Events	Total	Events	Total	Peto, Fixed, 95% CI		Peto, Fix	red, 95% CI	
Scognamiglio 1994	1	70	0	65	6.88 [0.14, 347.65]		. —	+	<u> </u>
						0.01	0.1	1 1	0 100
							Eoverire digovin	Egygurg golgiur	m ahannal blaakara

2

### Figure 48: Onset of symptoms or progression of NYHA class at 6 years

	Digox	rin	Calcium channel b	Risk Ratio	Risk Ratio							
Study or Subgroup	Events	Total	Events	Total	M-H, Fixed, 95% CI			M-H, Fixe	ed, 95% C	I		
Scognamiglio 1994	17 70		6	65	2.63 [1.11, 6.26]						_	
						0.1	0.2	0.5	1 2	2	5	10
								Favours digoxin	Favours	calcium char	nel bl	lockers

3

# Figure 49: Evidence of HVD progression on imaging (worsening of disease severity) at 6 years

		Digox	in	Calcium channel b	olocker	Peto Odds Ratio	Peto Odds Ratio					
Study or S	ubgroup	Events	Total	Events	Total	Peto, Fixed, 95% CI		Peto, Fixe	ed, 95% CI			
Scognamig	lio 1994	5	70	0	65	7.30 [1.23, 43.33]		1	<b>-</b>			
							0.01	0.1	10	100		
								Favours digoxin	Favours calcium ch	annel blockers		

4

### Figure 50: Need for heart valve intervention at 6 years



5

Figure 51: Withdrawal due to adverse events at 6 years



6

### E.173 Primary mitral stenosis

8 No studies identified.

## E.1.4 Primary mitral regurgitation

### E.1.421 ACE inhibitors compared to placebo

Figure 52: All-cause mortality at 6-12 months

	ACE-inhib	oitors	Placel	oo		Risk Difference	Risk Difference	
Study or Subgroup	Events			Events Total		M-H, Fixed, 95% CI	CI M-H, Fixed, 95% CI	
Marcotte 1997	0	6	0	10	34.8%	0.00 [-0.23, 0.23]	·] —	
Wisenbaugh 1994	0	12	1	17	65.2%	-0.06 [-0.22, 0.11]	j <del>-</del>	
Total (95% CI)		18		27	100.0%	-0.04 [-0.18, 0.11]	1	
Total events	0		1					
Heterogeneity: Chi <sup>2</sup> = 0.17, df = 1 (P = 0.68); Test for overall effect: $Z$ = 0.51 (P = 0.61)				o o			-1 -0.5 0 0.5 Favours ACE-I Favours placebo	<b>⊣</b> 1

3

Figure 53: Cardiac mortality at 6-12 months

_	ACE-inhi	oitors	Placel	bo		Risk Difference	Risk Difference
Study or Subgroup	Events	Total	<b>Events</b>	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
Marcotte 1997	0	6	0	10	34.8%	0.00 [-0.23, 0.23]	<del></del>
Wisenbaugh 1994	0	12	1	17	65.2%	-0.06 [-0.22, 0.11]	-
Total (95% CI)		18		27	100.0%	-0.04 [-0.18, 0.11]	•
Total events	0		1				
Heterogeneity: Chi2 =	0.17, $df = 1$	(P = 0.68)	8); $I^2 = 0\%$	6			1 05 0 05 1
Test for overall effect:	Z = 0.51 (P	= 0.61)					-1 -0.5 0 0.5 1 Favours ACE-I Favours placebo

4

5 6

7

Figure 54: Quality of life (life quality index, 1-6, change score, high is good) at 6 months

	ACE-	ACE-inhibitors		Placebo			Mean Difference	Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	IV, Fixed, 95% CI	IV, Fixed, 95% CI
Marcotte 1997	0.2	0.73	6	0.4	0.95	10	-0.20 [-1.03, 0.63]	<del></del>
							•	-4 -2 0 2 4
								Favours placebo Favours ACE-I

MIDs used to assess imprecision were calculated by multiplying the median baseline SD across study arms (2.24) by 0.5 and were ±1.12.

Figure 55: Quality of life (life quality index, 1-6, change score, high is good) at 1 year

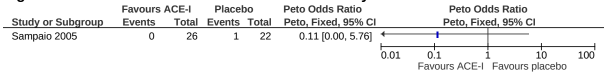
y ca.								
	ACE-	ACE-inhibitors			acebo		Mean Difference	Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	IV, Fixed, 95% CI	IV, Fixed, 95% CI
Marcotte 1997	0.3 0.73 6			0.4	0.95	10	-0.10 [-0.93, 0.73]	
							_	-4 -2 0 2 4
								Favours placeho Favours ACF-I

MIDs used to assess imprecision were calculated by multiplying the median baseline SD across study arms (2.24) by 0.5 and were  $\pm 1.12$ .

Figure 56: Onset of symptoms or progression of NYHA class at 6-12 months

	ACE-INNII	oitors	Placer	00		RISK Ratio		KIS	sk Ratio		
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI		M-H, F	ixed, 95%	CI	
Sampaio 2005	0	26	4	22	79.4%	0.09 [0.01, 1.67]	<b>←</b>				
Wisenbaugh 1994	0	12	1	17	20.6%	0.46 [0.02, 10.45]	_	-			
Total (95% CI)		38		39	100.0%	0.17 [0.02, 1.26]	-				
Total events	0		5								
Heterogeneity: Chi <sup>2</sup> = 0	).55, df = 1 (	P = 0.46	6); $I^2 = 0\%$	, D			0.01	0.1	+	10	100
Test for overall effect: 2	Z = 1.73 (P :	= 0.08)					0.01	Favours ACE	-l Favoui	s placebo	

Figure 57: Need for heart valve intervention at 1 year



2

Figure 58: Exercise tolerance (Bruce Protocol treadmill exercise time, seconds, change score) at 1 year

	ACE-	ACE-inhibitors		PI	acebo	Mean Difference			Mean Difference				
Study or Subgroup	Mean SD Total N		Mean	SD	Total	IV, Fixed, 95% CI		IV, Fix	ed, 95%	ed, 95% CI			
Marcotte 1997	39	61.2	6	18	66.4	10	21.00 [-42.97, 84.97]				1		
								-100	-50	Ó	50	100	
									Favours placebo	Favo	urs ACE-I		

MIDs used to assess imprecision were calculated by multiplying the median baseline SD across study arms (133.85) by 0.5 and were  $\pm 66.9$ .

5

# Figure 59: Exercise tolerance (oxygen uptake at peak exercise, mL/min, final value) at 1 year

•	ACE-inhibitors		Placebo			Mean Difference		ce				
Study or Subgroup	Mean SD Total		Mean	SD	Total	IV, Fixed, 95% CI		IV,	Fixed, 95%	G CI		
Sampaio 2005	1,794	561	26	1,433	521	21	361.00 [50.91, 671.09]				<del></del>	
							-1000	-500	Ó	500	1000	
									Favours A	CE-I Favo	urs placebo	ı

MIDs used to assess imprecision were calculated by multiplying the median baseline SD across study arms (541.0) by 0.5 and were ±270.5.

8

6 7

### Figure 60: Withdrawal due to adverse events at 1 year

	ACE-inhil	bitors	Place	bo	Risk Ratio		F	Risk Ratio		
Study or Subgroup	Events	Total	<b>Events</b>	Total	M-H, Fixed, 95% CI		M-H,	Fixed, 95%	6 CI	
Marcotte 1997	4	10	1	11	4.40 [0.59, 33.07]			+	<del>1</del> .	
						0.01	0.1	1	10	100

9

# E.1.402 Beta blockers compared to placebo

Figure 61: All-cause mortality at 2 years

	Beta-blo	cker	Place	bo	Peto Odds Ratio		Peto Oc	lds Ratio	
Study or Subgroup	Events	Total	<b>Events</b>	Total	Peto, Fixed, 95% CI		Peto, Fix	ed, 95% CI	
Ahmed 2012	1	19	0	18	7.01 [0.14, 353.80]		. ———	1.	
						0.01 0	.1	1 10	100
						Favours	beta-blocker	Favours placebo	

Figure 62: Cardiac mortality at 2 years

i igaic oz. c	ui aiuo		unty u	,	caio				
	Beta-blo	cker	Placel	00	Peto Odds Ratio		Peto Oc	lds Ratio	
Study or Subgroup	Events	Total	Events	Total	Peto, Fixed, 95% CI		Peto, Fix	ed, 95% CI	
Ahmed 2012	1	19	0	18	7.01 [0.14, 353.80]		. —	1.	
						0.01 (	).1 heta-blocker	1 10	100

Figure 63: Need for heart value intervention at 2 years



2

Figure 64: Serious adverse events at 2 years



3

# E.145 Primary tricuspid regurgitation

5 No studies identified.

# E.166 Secondary heart valve disease – mitral regurgitation or tricuspid regurgitation

7 No studies identified.

8

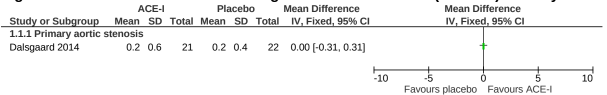
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### EL2 Valve disease with heart failure

### E.2.11 Primary aortic stenosis

#### E.2.1121 ACE-I versus placebo

Figure 65: Exercise tolerance: change in exercise duration (minutes) at 3 days



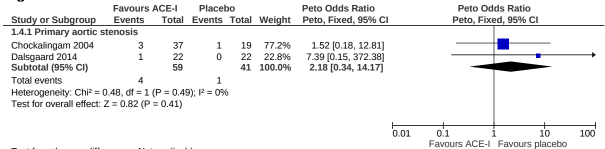
MIDs used to assess imprecision were calculated by multiplying the median baseline SD across study arms (2.0) by 0.5 and were ±1.0.

Figure 66: Exercise tolerance: 6-minute walk distance at 4 weeks

J	A	CE-I		Pla	acebo	)	Mean Difference		Mean D	ifference		
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	IV, Fixed, 95% CI		IV, Fixe	d, 95% CI		
1.2.1 Primary aortic	stenosis											
Chockalingam 2004	402	150	34	376	174	18	26.00 [-68.89, 120.89]			+ +		<b>→</b>
								-		-		——
								-100	-50	Ó	50	100
									Favours placebo	Favours	ACE-I	

MIDs used to assess imprecision were calculated by multiplying the median baseline SD across study arms (152.0) by 0.5 and were ±76.0.

Figure 67: Withdrawal due to adverse events at 2-3 months



Test for subgroup differences: Not applicable

1

### E.2.122 ARB versus placebo

#### Figure 68: All-cause mortality at 2-12 months

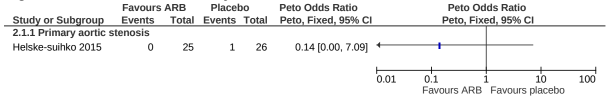


Figure 69: Incidence of acute heart failure at 2-12 months

_	Favours	ARB	Placel	bo	Peto Odds Ratio		Peto Od	lds Ratio	
Study or Subgroup	Events	Total	<b>Events</b>	Total	Peto, Fixed, 95% CI		Peto, Fix	ed, 95% CI	
2.2.1 Primary aortic s	stenosis								
Helske-suihko 2015	1	25	0	26	7.69 [0.15, 387.87]			1	<b></b>
						0.01	01	1 10	100
						0.01	Favours ARB	Favours placebo	100

Figure 70: Exercise tolerance: change from baseline 6-minute walking distance at 2-12 months

	A	ARB		Pla	aceb	0	Mean Difference		Mean Difference	ce	
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	IV, Fixed, 95% Cl		IV, Fixed, 95%	CI	
2.3.1 Primary aortic s	stenosis										
Helske-suihko 2015	-20	42	22	-2	59	21	-18.00 [-48.74, 12.74]		<del>- + -</del>		
								-100	-50 0	50	100
									Favours placebo Favou	ırs ARB	

MIDs used to assess imprecision were calculated by multiplying the median baseline SD across study arms (148) by 0.5 and were ±74.0.

5

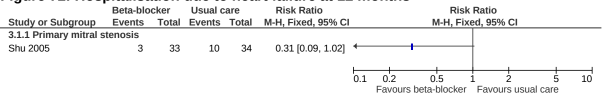
Figure 71: Withdrawal due to adverse events at 2-12 months

	ARE	3	Placel	bo	Risk Ratio			Risk	Ratio			
Study or Subgroup	Events	Total	<b>Events</b>	Total	M-H, Fixed, 95% CI			M-H, Fixe	ed, 95%	CI		
2.4.1 Primary aortic s	tenosis											
Helske-suihko 2015	2	25	2	26	1.04 [0.16, 6.83]						_	
						0.1	0.2	0.5	Favour	2 5	1	0

## E.2.2 Primary mitral stenosis

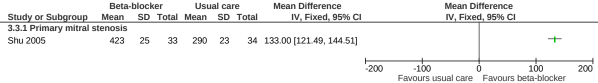
#### E.2.221 Beta-blocker versus usual care

Figure 72: Hospitalisation due to heart failure at 12 months



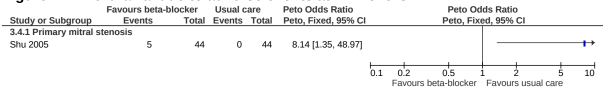
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Figure 73: Exercise tolerance: 6-minute walking distance at 6-12 months



MIDs used to assess imprecision were calculated by multiplying the median baseline SD across study arms (30.0) by 0.5 and were  $\pm 15.0$ .

Figure 74: Withdrawal due to adverse events at 12 months



4

### E.2.252 Beta-blocker versus placebo

Figure 75: Exercise tolerance: treadmill exercise time to exhaustion at 1-4 weeks

	Favou	rs plac	ebo	Pla	acebo	1		Mean Difference		Mean Difference	
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Random, 95% CI		IV, Random, 95% CI	
4.1.1 Primary mitral	stenosis										
Bassan 1987	4.57	1.32	10	4.72	1.37	10	34.7%	-0.15 [-1.33, 1.03]		<b></b>	
Klein 1985	11	2	13	9	2	13	29.7%	2.00 [0.46, 3.54]		<del></del>	
Patel 1995 Subtotal (95% CI)	8.8	1.7	19 <b>42</b>	9.4	1.8	19 <b>42</b>	35.6% <b>100.0</b> %	-0.60 [-1.71, 0.51] <b>0.33 [-1.09</b> , 1.75]			
Heterogeneity: Tau² = Test for overall effect:				(P = 0.0	02); I²	= 74%					
									-10	-5 0 5 Favours placebo Favours beta-blocke	10 r

Test for subgroup differences: Not applicable

MIDs used to assess imprecision were calculated by multiplying the median final value control group SD across studies (1.8) by 0.5 and were ±0.9.

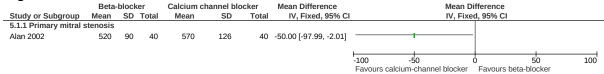
# Figure 76: Exercise tolerance: pulmonary capillary wedge pressure after exercise at 6 months

	Beta-	block	cer	PI	acebo	)	Mean Difference		Mean Di	ifference	
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	IV, Fixed, 95% CI		IV, Fixe	d, 95% CI	
4.2.1 Primary mitral	stenosis										
Kumar 1994	35.7	7.3	13	50.5	10.4	13	-14.80 [-21.71, -7.89]		-		
								-50	-25	0 25	50
								Fa	vours beta-blocker	Favours placebo	

MIDs used to assess imprecision were calculated by multiplying the median baseline SD across study arms (10.7) by 0.5 and were ±5.35.

#### E.2.243 Beta-blocker versus calcium channel blocker

#### Figure 77: Exercise tolerance: total effort time on treadmill exercise test at 3 months



MIDs used to assess imprecision were calculated by multiplying the median baseline SD across study arms (120) by 0.5 and were ±60.0.

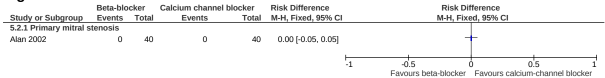
Note: baseline total effort time not matched - beta-blocker: 452±120; calcium-channel blocker: 534±120

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Figure 78: Withdrawal due to adverse events



6

# E.273 Secondary heart valve disease (mitral regurgitation and tricuspid regurgitation)

#### Figure 79: Cardiac mortality at 12 weeks

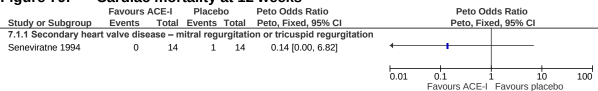
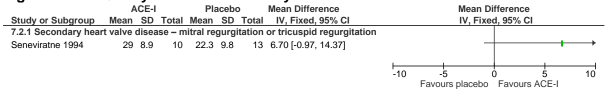


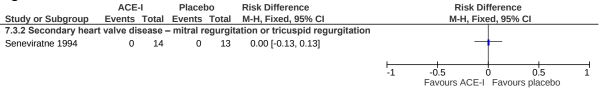
Figure 80: Quality of life: Duke activity index score at 12 weeks



MIDs used to assess imprecision were calculated by multiplying the median baseline SD across study arms (9.4) by 0.5 and were ±4.7.

1

### Figure 81: Withdrawal due to adverse events at 3 months



# Appendix F: GRADE tables

F.3 Valve disease without heart failure

# F.14 Primary aortic [including bicuspid] stenosis

5 Table 33: Clinical evidence profile: ACE inhibitors compared to placebo

			Quality as	sessment			No of patients Effect  ACE- Relative ALL LANGE		Effect	Quality	Importance	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	ACE- inhibitors	Placebo	Relative (95% CI)	Absolute		
All-cause	mortality - no	t reported										
0	=	_	=	-	_	none	-	-	-	-		CRITICAL
Cardiac m	ortality - not	reported										
0	-	-	-	-	-	none	-	-	-	-		CRITICAL
Health-rela	ated quality o	f life - not	reported									
0	-	-	-	-	-	none	-	-	-	-		CRITICAL
Health-rela	ated quality o	f life - not	reported									
0	-	-	-	-	-	none	-	-	-	-		CRITICAL
Onset of s	symptoms or	progression	on in NYHA class	not reported								
0	-	-	-	-	-	none	-	-	-	-		CRITICAL
Evidence (	of HVD progre	ession on	imaging (worseni	ng of disease se	verity) - not rep	orted						
0	-	-	-	-	-	none	-	-	-	-		CRITICAL
Need for h	eart valve int	ervention	(follow-up 12 mor	nths)								
1		- ,	no serious inconsistency	no serious indirectness	very serious <sup>2</sup>	none	4/40 (10%)	4.7%	RR 2.15 (0.42 to 11.1)	54 more per 1000 (from 27 fewer to 475 more)	⊕OOO VERY LOW	CRITICAL
Exercise to	olerance (cha	inge score	e) (follow-up 12 mo	onths; measured	with: Exercise	distance measured	d with tread	nill exerc	ise test; Bette	er indicated by lower va	lues)	
		,	no serious inconsistency		no serious imprecision³	none	26	41	-	MD 49 lower (61.59 to 36.41 lower)	⊕⊕OO LOW	IMPORTANT
Withdrawa	al due to adve	rse event	s (follow-up 12 mo	onths)								

				no serious indirectness	very serious <sup>2</sup>	none	2/38 (5.3%)	2.4%	RR 2.21 (0.21 to 23.41)	29 more per 1000 (from 19 fewer to 538 more)	⊕OOO VERY LOW	IMPORTANT
--	--	--	--	----------------------------	---------------------------	------	----------------	------	----------------------------	---	---------------------	-----------

<sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias <sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

Table 34: Clinical evidence profile: beta blockers compared to placebo

			Quality ages	.coment			No of no	ntionto		Effect		
			Quality asse	ssment			No of pa	atients		Enect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Beta- blockers	Placebo	Relative (95% CI)	Absolute		
All-cause	mortality - not	reported										
0	-	-	-	-	-	none	-	-	-	-		CRITICAL
Cardiac m	ortality - not r	eported										
0	-	-	-	-	_	none	-	-	ı	-		CRITICAL
Health-rela values)	ated quality of	f life (chan	ge score) (follow-u	ip 5 months; mea		Minnesota living w	ith heart fa	ilure que	stionnaire; ra	ange of scores: 0-105; Bet	ter indica	
	randomised trials	very serious¹	no serious inconsistency	no serious indirectness	serious <sup>2,3</sup>	none	19	19	1	MD 6 higher (0.55 lower to 12.55 higher)	⊕OOO VERY LOW	CRITICAL
Health-rela	ated quality of	f life - not ı	reported									
0	-	-	=	-	-	none	-	-	-	-		CRITICAL
Onset of s	ymptoms or p	orogressio	n in NYHA class -	not reported								
0	-	-	=	-	-	none	-	-	-	-		CRITICAL
Evidence	of HVD progre	ession on i	maging (worsening	g of disease seve	erity) - not re	ported						
0	-	-	=	-	-	none	-	-	-	-		CRITICAL
Need for h	eart valve into	ervention -	not reported									
0	-	-	=	-	-	none	-	-	-	-		CRITICAL
Exercise t	olerance (cha	nge score	(follow-up 5 mont	hs; measured wi	th: 6 minute	walk test distance	Better indi	icated by	lower values	s)		
	randomised trials	very serious¹	no serious inconsistency	no serious indirectness	serious <sup>2,4</sup>	none	19	19	-	MD 12 lower (42.22 lower to 18.22 higher)	⊕OOO VERY LOW	IMPORTANT
Withdrawa	al or dose red	uction due	to adverse events	(follow-up 5 mo	nths)							
	randomised trials		no serious inconsistency	serious <sup>5</sup>	very serious²	none	4/19 (21.1%)	10.5%	RR 2 (0.41 to 9.65)	105 more per 1000 (from 62 fewer to 908 more)	⊕000 VERY LOW	IMPORTANT

<sup>&</sup>lt;sup>3</sup> MIDs used to assess imprecision were ±187.0

6

- <sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias <sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs
- <sup>3</sup> MIDs used to assess imprecision were ±5.0
- <sup>4</sup> MIDs used to assess imprecision were 21.0
- <sup>5</sup> Downgraded by 1 increment as the outcome includes people who had dose reductions or withdrawal due to adverse events

Table 35: Clinical evidence profile: diuretics compared to placebo

			Quality ass	essment			No of patients  Diuretics Placeb			Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Diuretics	Placebo	Relative (95% CI)	Absolute		
All-cause	mortality (follo	ow-up 19 ı	months)									
1	randomised trials		no serious inconsistency	no serious indirectness	very serious <sup>2</sup>	none	1/30 (3.3%)	6.5%	RR 0.52 (0.05 to 5.4)	31 fewer per 1000 (from 62 fewer to 286 more)	⊕OOO VERY LOW	CRITICAL
Cardiac m	ortality (follow	v-up 19 m	onths)									
1	randomised trials		no serious inconsistency	no serious indirectness	very serious <sup>2</sup>	none	0/29 (0%)	3.3%	OR 0.14 (0 to 7.06)	30 fewer per 1000 (from 120 fewer to 60 more) <sup>3</sup>	⊕OOO VERY LOW	CRITICAL
Health-rel	ated quality of	f life - not	reported									
0	-	-	-	-	1	none	-	-	-	-		CRITICAL
Health-rel	ated quality of	life (char	ge score) (follow-	up 12 months; m	easured with:	SF-36 physical fu	nctioning	subsca	e; range of sco	res: 0-100; Better indicate	d by high	ner values)
1	randomised trials		no serious inconsistency	no serious indirectness	very serious <sup>2,4</sup>	none	29	30	-	MD 4 higher (6.5 lower to 14.5 higher)	⊕OOO VERY LOW	CRITICAL
Health-rel	ated quality of	life (char	ge score) (follow-	up 12 months; m	easured with:	SF-36 role physic	al subsca	le; rang	e of scores: 0-1	00; Better indicated by high	gher value	
1	randomised trials		no serious inconsistency	no serious indirectness	very serious <sup>2,4</sup>	none	29	30	-	MD 3 higher (15.12 lower to 21.12 higher)	⊕OOO VERY LOW	CRITICAL
Onset of s	symptoms or p	progression	n of NYHA class (	follow-up 19 mor	nths)							
1	randomised trials		no serious inconsistency	no serious indirectness	very serious <sup>2</sup>	none	13/29 (44.8%)	33.3%	RR 1.34 (0.7 to 2.57)	113 more per 1000 (from 100 fewer to 523 more)	⊕OOO VERY LOW	CRITICAL
Evidence	of HVD progre	ession on	imaging (worsenir	ng of disease sev	erity) - not re	ported						
0	-	-	-	-	-	none	-	-	-	-		CRITICAL
Need for h	eart valve into	ervention	not reported									
0	-	-	-	-	-	none	-	-	-	-		CRITICAL
Withdrawa	al due to adve	rse events	(follow-up 19 mo	nths)								

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1	randomised trials	serious <sup>1</sup>	no serious inconsistency	no serious indirectness	very serious <sup>2</sup>	none	1/32 (3.1%)	0%	OR 6.94 (0.14 to 350.54)	30 more per 1000 (from 50 fewer to 120 more) <sup>3</sup>	⊕000 VERY LOW	IMPORTANT
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<sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias <sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs <sup>3</sup> Absolute effect calculated manually using risk difference as zero events in one arm of the study <sup>4</sup> MIDs used to assess imprecision were ±3.0

Table 36: Clinical evidence profile: statins compared to placebo 5

			Quality asse	Quality assessment  consistency Indirectness Imprecision Other						Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Statins	Placebo	Relative (95% CI)	Absolute		
All-cause	mortality (foll	low-up mean	4.3 years)									
-	randomised trials	serious <sup>1</sup>	no serious inconsistency	serious <sup>2</sup>	very serious <sup>3</sup>	none	109/1102 (9.9%)	4.4%	RR 1.01 (0.79 to 1.3)	0 more per 1000 (from 9 fewer to 13 more)	⊕OOO VERY LOW	CRITICAL
All-cause	mortality (tim	e to event) (f	ollow-up 4.4 years	s)								
	randomised trials	serious <sup>1</sup>	no serious inconsistency	serious <sup>2</sup>	very serious <sup>3</sup>	none	105/944 (11.1%)		HR 1.04 (0.79 to 1.37)	4 more per 1000 (from 22 fewer to 37 more)	⊕OOO VERY LOW	CRITICAL
Cardiac m	ortality (follo	w-up mean 3	.7 years)		•			•				
			no serious inconsistency	serious <sup>2</sup>	serious <sup>3</sup>	none	61/1179 (5.2%)	5.2%	RR 0.75 (0.54 to 1.03)	13 fewer per 1000 (from 24 fewer to 2 more)	⊕⊕OO LOW	CRITICAL
Cardiac m	ortality (time	to event) (fol	low-up 4.4 years)	<b>'</b>			ļ.	!	<u> </u>			
	randomised trials	serious <sup>1</sup>	no serious inconsistency	serious <sup>2</sup>	very serious <sup>3</sup>	none	47/944 (5%)	56/929 (6%)	HR 0.83 (0.56 to 1.23)	10 fewer per 1000 (from 26 fewer to 13 more)	⊕OOO VERY LOW	CRITICAL
Health-rel	ated quality o	f life - not rep	oorted				*	•				
0	-	_	-	-	-	none	-	-	-	-		CRITICAL
Health-rel	ated quality o	f life - not rep	oorted									
0	-	-	-	-	-	none	-	-	-	-		CRITICAL
Onset of s	symptoms or	progression	of NYHA class (fo									
			no serious inconsistency	serious <sup>2</sup>	very serious <sup>3</sup>	none	28/1021 (2.7%)	4.4%	RR 0.99 (0.59 to 1.66)	0 fewer per 1000 (from 18 fewer to 29 more)	⊕OOO VERY LOW	CRITICAL
Onset of s	symptoms or	progression	of NYHA class (tin	ne to event) (foll	ow-up 4.4 years	)		•				

All rights roson

# F.162 Primary aortic regurgitation

7 Table 37: Clinical evidence profile: ACE inhibitors compared to placebo/no treatment

			Quality as	sessment			No of	patients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	ACE- inhibitors	Placebo/no treatment	Relative (95% CI)	Absolute		
All-cause	mortality (fo	llow-up 7	years)									
1	randomised trials			no serious indirectness	very serious <sup>2</sup>	none	1/32 (3.1%)	3.2%	RR 0.97 (0.06 to 14.82)	1 fewer per 1000 (from 30 fewer to 442 more)	⊕OOO VERY LOW	CRITICAL
Cardiac n	nortality (follo	ow-up 7 ye	ears)		•	•	,				•	

<sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

<sup>&</sup>lt;sup>2</sup> Downgraded by 1 increment as one study included a statin and ezetimibe in the intervention group

<sup>3</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

<sup>&</sup>lt;sup>4</sup> Downgraded by 1 as the point estimate varies widely across studies, with subgroup analysis not being possible due to the difference being seen in one study

<sup>&</sup>lt;sup>5</sup> Absolute effect calculated manually using risk difference as zero events in one arm of the study

1	randomised trials	very serious <sup>1</sup>	no serious inconsistency	no serious indirectness	very serious <sup>2</sup>	none	0/32 (0%)	3.2%	OR 0.13 (0 to 6.61)	30 fewer per 1000 (from 120 fewer to 50 more) <sup>3</sup>	⊕OOO VERY LOW	CRITICAL
Health-re	lated quality	of life - no	t reported									
0	-	=	_	-	-	none	-	-	-	-		CRITICAL
Health-re	lated quality	of life - no	t reported									
0	=	-	-	=	=	none	-	-	-	-		CRITICAL
Onset of	symptoms or	progress	ion of NYHA clas	s (follow-up 7 y	ears)							
2	randomised trials	serious <sup>1</sup>	serious <sup>4</sup>	no serious indirectness	no serious imprecision	none	10/43 (23.3%)	8/40 (20%)	RD 0 (-0.13 to 0.22)	40 more per 1000 (from 130 fewer to 220 more) <sup>5</sup>	⊕⊕OO LOW	CRITICAL
Evidence	of HVD prog	ression o	n imaging (worse	ning of disease	severity) (follow	w-up 7 years)						
1	randomised trials	very serious¹	no serious inconsistency	no serious indirectness	very serious <sup>2</sup>	none	14/32 (43.8%)	32.3%	RR 1.36 (0.71 to 2.58)	116 more per 1000 (from 94 fewer to 510 more)	⊕000 VERY LOW	CRITICAL
Need for	heart valve in	terventio	n (follow-up 7 yea	ars)								
1	randomised trials	very serious <sup>1</sup>	no serious inconsistency	no serious indirectness	very serious <sup>2</sup>	none	16/32 (50%)	38.7%	RR 1.29 (0.74 to 2.27)	112 more per 1000 (from 101 fewer to 491 more)	⊕000 VERY LOW	CRITICAL
Withdraw	al due to adv	erse even	nts (follow-up 7 ye	ears)								
1	randomised trials	very serious¹	no serious inconsistency	no serious indirectness	very serious <sup>2</sup>	none	3/32 (9.4%)	0%	OR 7.65 (0.77 to 76.34)	90 more per 1000 (from 20 fewer to 210 more) <sup>3</sup>	⊕OOO VERY LOW	IMPORTANT

<sup>&</sup>lt;sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias <sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

# Table 38: Clinical evidence profile: ACE inhibitors compared to calcium channel blockers

			Quality as:	sessment			No of	patients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	ACE- inhibitors	Calcium channel blockers	Relative (95% CI)	Absolute		
All-cause	mortality (fo	llow-up 7	years)				·					
1	randomised trials	,		no serious indirectness	very serious <sup>1</sup>	none	1/32 (3.1%)	1/32 (3.1%)		0 fewer per 1000 (from 29 fewer to 447 more)		CRITICAL

Absolute effect calculated manually using risk difference as zero events in one arm of the study
 Downgraded by 1 increment as zero events in both arms of one study
 Absolute effect calculated manually using risk difference as zero events in both arms of a study

**CRITICAL** 

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0/32

(0%)

1/32

OR 0.14 (0 to

6.82)

30 fewer per 1000

(from 110 fewer to 50

no serious

inconsistency

no serious

indirectness

verv serious1

none

Cardiac mortality (follow-up 7 years) randomised

trials

verv

serious<sup>2</sup>

# Table 39: Clinical evidence profile: ARBs compared to beta blockers

			Quality assessn	nent			No	of patients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	ARB	KOTA-	Relative (95% CI)	Absolute		
All-cause m	nortality - not re	ported			•							•
0	-	-	-	-	-	none	-	-	-	-		CRITICAL

<sup>&</sup>lt;sup>2</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

<sup>&</sup>lt;sup>3</sup> Absolute effect calculated manually using risk difference as zero events in one arm of the study

<sup>&</sup>lt;sup>4</sup> Absolute effect calculated manually using risk difference as zero events in both arms of one study

<sup>&</sup>lt;sup>5</sup> Downgraded by 1 increment as zero events in one of the studies included

<sup>&</sup>lt;sup>6</sup> Absolute effect calculated manually using risk difference as zero events in one arm of a study

Cardiac mo	rtality - not rep	orted										
0	=	-	-	-	_	none	-	-	-	-		CRITICAL
Health-relat	ted quality of lif	e - not repo	orted					·				
0	-	-	-	-	_	none	-	-	-	-		CRITICAL
Health-relat	ted quality of lif	e - not repo	orted									
0	-	-	-	-	_	none	-	-	-	-		CRITICAL
Onset of sy	mptoms or pro	gression in	NYHA class - not re	ported								
0	-	=	-	-	_	none	-	=	-	-		CRITICAL
Evidence of	f HVD progress	ion on ima	ging (worsening of d	isease severi	ty) - not repo	rted						
0	-	=	-	-	_	none	-	=	-	-		CRITICAL
Need for he	art valve interv	ention - no	t reported									
0	-	-	-	-	_	none	-	=	-	-		CRITICAL
Exercise to	lerance (final v	alue) (follov	v-up 3 weeks; measu	red with: exe	rcise work ra	ate using an ergome	ter; B	etter indica	ted by hi	gher values)		
	randomised trials		no serious inconsistency		very serious <sup>3,4</sup>	none	17	17	-	MD 0 higher (4.75 lower to 4.75 higher)	⊕OOO VERY LOW	IMPORTANT

<sup>&</sup>lt;sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias <sup>2</sup> Downgraded by 1 increment as follow up less than 1 month <sup>3</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

Table 40: Clinical evidence profile: beta blockers compared to placebo

			Quality asses	ssment			No of pa	ntients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Beta- blocker	Placebo	Relative (95% CI)	Absolute		
All-cause n	nortality - not r	eported						•				
0	-	-	-	-	_	none	-	-	-	=		CRITICAL
Cardiac mo	ortality - not re	ported										
0	-	-	-	-	-	none	-	-	-	-		CRITICAL
Onset of sy	mptoms or pr	ogression	in NYHA class - not	reported								
0	-	=	-	-	-	none	-	-	-	-		CRITICAL
Quality of I	ife (final value)	(follow-up	6 months; measur	ed with: EuroQol v	isual analog	ue scale; range of s	scores: 0-1	00; Bette	r indicate	ed by higher values)		
	randomised trials	serious <sup>1</sup>		no serious indirectness	serious <sup>2,3</sup>	none	36	36	-	MD 3 higher (2.7 lower to 8.7 higher)	⊕⊕OO LOW	CRITICAL
Quality of I	ife (final value)	(follow-up	6 months; measure	ed with: KCCQ; rai	nge of score	s: 0-100; Better indi	cated by h	igher val	ues)			

<sup>&</sup>lt;sup>4</sup> MIDs used to assess imprecision were ±4.0

1	randomised trials	serious <sup>1</sup>		no serious indirectness	serious <sup>2,4</sup>	none	36	36	-	MD 2 higher (17.76 lower to 21.76 higher)	⊕⊕OO LOW	CRITICAL
Evidence o	of HVD progres	sion on in	naging (worsening o	f disease severity)	- not report	ed						
0	-	-	-	-	-	none	-	-	-	-		CRITICAL
Need for h	eart valve inter	vention - r	not reported									
0	-	-	-	-	-	none	-	-	-	-		CRITICAL
Exercise to	olerance (follow	v-up 6 moi	nths; measured with	: Peak work (bicyo	le ergomete	r); Better indicated I	oy higher v	alues)				
1		serious <sup>1</sup>			serious <sup>2,5</sup>	none	36	36	-	MD 12 lower (40.64 lower to		IMPORTANT
	trials		inconsistency	indirectness						16.64 higher)	LOW	

<sup>&</sup>lt;sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias <sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs <sup>3</sup> MIDs used to assess imprecision were ±5.00 <sup>4</sup> MIDs used to assess imprecision were ±21.39 <sup>5</sup> MIDs used to assess imprecision were ±31.50

Table 41: Clinical evidence profile: calcium channel blockers compared to placebo/no treatment

			Quality asso	essment			No of p	patients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Calcium channel blockers	Placebo/no treatment	Relative (95% CI)	Absolute		
All-cause	mortality (fo	llow-up 7	years)									
1	randomised trials			no serious indirectness	very serious²	none	1/32 (3.1%)	3.2%	RR 0.97 (0.06 to 14.82)	1 fewer per 1000 (from 30 fewer to 442 more)	⊕OOO VERY LOW	CRITICAL
Cardiac n	nortality (follo	ow-up 7 ye	ears)									
1	randomised trials			no serious indirectness	very serious <sup>2</sup>	none	1/32 (3.1%)	3.1%	RR 1 (0.07 to 15.3)	0 fewer per 1000 (from 29 fewer to 443 more)	⊕OOO VERY LOW	CRITICAL
Health-re	lated quality	of life - no	ot reported									
0	-	-	-	-	-	none	-	-	-	-		CRITICAL
Health-re	ated quality	of life - no	ot reported									
0	-	-	-	-	-	none	-	-	-	-		CRITICAL
Onset of	symptoms or	progress	ion of NYHA clas	s (follow-up 7 ye	ears)				•		•	
1	randomised trials	- ,		no serious indirectness	very serious <sup>2</sup>	none	8/32 (25%)	25.8%	RR 0.97 (0.42 to 2.26)	8 fewer per 1000 (from 150 fewer to 325 more)	⊕OOO VERY LOW	CRITICAL

Evidence	Evidence of HVD progression on imaging (worsening of disease severity) (follow-up 7 years)													
1		- , .		no serious indirectness	very serious <sup>2</sup>	none	10/32 (31.3%)	32.3%	RR 0.97 (0.47 to 2)	10 fewer per 1000 (from 171 fewer to 323 more)	⊕OOO VERY LOW	CRITICAL		
Need for	Need for heart valve intervention (follow-up 7 years)													
2	randomised trials	serious <sup>1</sup>		no serious indirectness	very serious <sup>4</sup>	none	13/68 (19.1%)	12/67 (17.9%)	RD 0.01 (- 0.11 to 0.13)	10 more per 1000 (from 110 fewer to 130 more) <sup>5</sup>	⊕OOO VERY LOW	CRITICAL		
Withdraw	val due to adv	erse ever	nts (follow-up 7 ye	ars)										
2	randomised trials	very serious¹		no serious indirectness	serious <sup>2</sup>	none	8/70 (11.4%)	0%	RR 8.51 (1.12 to 64.44)	120 more per 1000 (from 30 more to 200 more) <sup>6</sup>	⊕000 VERY LOW	IMPORTANT		

<sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

# Table 42: Clinical evidence profile: digoxin compared to calcium channel blockers

			Quality asse	essment			No	of patients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Digoxin	Calcium channel blockers	Relative (95% CI)	Absolute		
All-cause	mortality (foll	ow-up 6 y	/ears)									
1	randomised trials	serious <sup>1</sup>	no serious inconsistency	no serious indirectness	very serious²	none	1/70 (1.4%)	0/65 (0%)	OR 6.88 (0.14 to 347.65)	10 more per 1000 (from 30 fewer to 50 more) <sup>3</sup>	⊕OOO VERY LOW	CRITICAL
Cardiac m	nortality - not	reported	,	<u>-                                    </u>								
0	-	_	-	-	-	none	-	=	-	=		CRITICAL
Health-rel	ated quality o	f life - not	reported									
0	-	_	-	_	-	none	=	-	-	-		CRITICAL
Health-rel	ated quality o	f life - not	reported									
0	_			-	-	none	-	-	-	-		CRITICAL
Onset of	symptoms of p	orogressi	on of NYHA class	(follow-up 6 yea	rs)							
1	randomised trials	serious <sup>1</sup>	no serious inconsistency	no serious indirectness	serious <sup>2</sup>	none	17/70 (24.3%)	6/65 (9.2%)	RR 2.63 (1.11 to 6.26)	150 more per 1000 (from 10 more to 486 more)	⊕⊕OO LOW	CRITICAL

<sup>&</sup>lt;sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

Downgraded by 1 increment as zero events in both arms of one study
 Downgraded by 2 increments as zero events in both arms of a study and OIS <80%</li>

Absolute effect calculated manually using risk difference as zero events in both arms of a study
 Absolute effect calculated manually using risk difference as zero events one arm of the study

Evidence	Evidence of HVD progression on imaging (worsening of disease severity) (follow-up 6 years)													
1	randomised trials		no serious inconsistency	no serious indirectness	serious <sup>2</sup>	none	5/70 (7.1%)	0/65 (0%)		70 fewer per 1000 (from 10 more to 140 more) <sup>3</sup>	⊕⊕OO LOW	CRITICAL		
Need for I	leed for heart valve intervention (follow-up 6 years)													
1	randomised trials		no serious inconsistency	no serious indirectness	serious <sup>2</sup>	none	20/70 (28.6%)	6/65 (9.2%)	RR 3.10 (1.33 to 7.22)	194 more per 1000 (from 30 more to 574 more)	⊕⊕OO LOW	CRITICAL		
Withdraw	Withdrawal due to adverse events (follow-up 6 years)													
1	randomised trials		no serious inconsistency	no serious indirectness	serious <sup>4</sup>	none	0/70 (0%)	0/65 (0%)	RD 0 (-0.03 to 0.03)	0 fewer per 1000 (from 30 fewer to 30 more) <sup>5</sup>	⊕⊕OO LOW	IMPORTANT		

<sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

# **Primary mitral stenosis**

No studies identified.

# **Primary mitral regurgitation**

Table 43: Clinical evidence profile: ACE inhibitors compared to placebo 9

	Design   Inconsistency   Indirectness   Imprecision									Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	ACE- inhibitors	Placebo	bo Relative (95% CI) Absolute			
All-cause	mortality (fol	low-up 6-1	12 months)									
	randomised trials	very serious <sup>1</sup>	serious <sup>2</sup>	serious <sup>3</sup>	very serious <sup>4</sup>	none	0/18 (0%)	2.9%		40 fewer per 1000 (from 180 fewer to 110 more) <sup>5</sup>	⊕OOO VERY LOW	CRITICAL
Cardiac m	ortality (follo	w-up 6-12	months)									
	randomised trials	very serious <sup>1</sup>	serious <sup>2</sup>	serious <sup>3</sup>	very serious <sup>4</sup>	none	0/18 (0%)	2.9%		40 fewer per 1000 (from 180 fewer to 110 more) <sup>5</sup>	⊕OOO VERY LOW	CRITICAL

<sup>&</sup>lt;sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

<sup>&</sup>lt;sup>3</sup> Absolute effect calculated from risk difference due to zero events in one study arm

<sup>&</sup>lt;sup>4</sup> Downgraded by 1 increment as sample size is between 75 and 350 with zero events in both arms <sup>5</sup> Absolute effect calculated from risk difference due to zero events in both study arms

					,				1			
1	randomised	very	no serious	serious <sup>3</sup>	no serious	none	6	10	-	MD 0.2 lower (1.03	$\oplus$ OOO	CRITICAL
	trials	serious1	inconsistency		imprecision <sup>6</sup>					lower to 0.63 higher)	VERY	
											LOW	
Quality o	f life (change	score) (fo	llow-up 1 years; n	neasured with: Li	ife quality index	; range of scores:	1-6; Better ir	ndicated	by higher valu	ies)		
1	randomised	very	no serious	serious <sup>3</sup>	no serious	none	6	10	-	MD 0.1 lower (0.93	$\oplus$ OOO	CRITICAL
	trials	serious1	inconsistency		imprecision <sup>6</sup>					lower to 0.73 higher)	VERY	
											LOW	
Onset of	symptoms or	progress	ion of NYHA class	(follow-up 6-12	months)				•			
2	randomised	very	no serious	serious <sup>3</sup>	very serious <sup>7</sup>	none	0/38	12%	RR 0.17 (0.02	140 fewer per 1000	⊕ООО	CRITICAL
	trials	serious1	inconsistency				(0%)		to 1.26)	(from 270 fewer to 10	VERY	
										fewer)⁵	LOW	
Evidence	of HVD progr	ession or	n imaging (worsen	ing of disease se	everity) - not rep	orted			•			
0	-	-	=	=	=	none	-	=	-	=		CRITICAL
Need for	heart valve in	tervention	n (follow-up 1 year	s)		•			•			
1	randomised	very	no serious	no serious	very serious <sup>7</sup>	none	0/26	4.6%	OR 0.11 (0 to	50 fewer per 1000 (from	⊕000	CRITICAL
	trials	serious1	inconsistency	indirectness			(0%)		5.76)	160 fewer to 70 more) <sup>5</sup>	VERY	
										·	LOW	
Exercise	tolerance (cha	ange scor	e) (follow-up 1 year	ars, measured w	ith: Bruce Proto	col treadmill exerc	ise time; Be	tter indi	cated by lower	values)		
1	randomised	very	no serious	serious <sup>3</sup>	serious <sup>7,8</sup>	none	6	10	-	MD 21 higher (42.97	⊕OOO	IMPORTANT
	trials	serious1	inconsistency							lower to 84.97 higher)	VERY	
			·							• ,	LOW	
Exercise	tolerance (fina	al value) (	follow-up 1 years;	measured with:	oxygen uptake	at peak exercise; E	Better indica	ted by h	igher values)			
1	randomised	very	no serious	no serious	serious <sup>7,9</sup>	none	26	21	-	MD 361 higher (50.91 to	⊕ООО	IMPORTANT
	trials	serious1	inconsistency	indirectness						671.09 higher)	VERY	
											LOW	
Withdrav	val due to adve	erse even	ts (follow-up 1 yea	ars)	•	•						
1	randomised	serious <sup>1</sup>	no serious	serious <sup>3</sup>	very serious <sup>7</sup>	none	4/10	9.1%	RR 4.4 (0.59	309 more per 1000	⊕000	IMPORTANT
	trials		inconsistency				(40%)		to 33.07)	(from 37 fewer to 1000	VERY	
							( , . ,				V L I \ I	

Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias powngraded by 1 increment as one study has zero events in both arms, and one has zero events in one arm Downgraded by 1 increment as some of the participants in one study may have had congenital valvular heart disease Downgraded by 2 increments as calculated power was less than 80%

Absolute effect calculated manually using risk difference as zero events in the studies

<sup>&</sup>lt;sup>6</sup> MIDs used to assess imprecision were ±1.12

4 5

Table 44: Clinical evidence profile: beta blockers compared to placebo

1 4.5.0		o v i a o i i	ce prome. be	ia bioonoi o	omparoa	to placebo						
		Quality asse	essment			No of p	atients		Effect	Quality	Importance	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Beta- blocker	Placebo	Relative (95% CI)	Absolute		
All-cause	mortality (follo	ow-up 2 ye	ears)									
1	randomised trials	serious <sup>1</sup>	no serious inconsistency	no serious indirectness	very serious²	none	1/19 (5.3%)	0%	OR 7.01 (0.14 to 353.8)	50 more per 1000 (from 80 fewer to 190 more) <sup>3</sup>	⊕000 VERY LOW	CRITICAL
Cardiac m	ortality (follow	v-up 2 yea	ırs)									
1	randomised trials	serious <sup>1</sup>	no serious inconsistency	no serious indirectness	very serious²	none	1/19 (5.3%)	0%	OR 7.01 (0.14 to 353.8)	50 more per 1000 (from 80 fewer to 190 more) <sup>3</sup>	⊕OOO VERY LOW	CRITICAL
Health-rela	ated quality of	f life - not	reported									
0	=	-	-	-	-	none	=	-	-	-		CRITICAL
Health-rela	ated quality of	f life - not	reported									
0	1	-	-	-	_	none	-	-	-	-		CRITICAL
Onset of s	ymptoms or p	orogressio	on in NYHA class -	not reported								
0	-	-	-	-	-	none	-	-	-	-		CRITICAL
Evidence	of HVD progre	ession on	imaging (worsenir	ng of disease sev	erity) - not re	ported						
0	-	-	-	-	-	none	-	-	_	-		CRITICAL
Need for h	eart valve into	ervention	(follow-up 2 years	)								
1	randomised trials	serious <sup>1</sup>	no serious inconsistency	no serious indirectness	very serious²	none	2/18 (11.1%)	33.3%	RR 0.33 (0.08 to 1.44)	223 fewer per 1000 (from 306 fewer to 147 more)	⊕OOO VERY LOW	CRITICAL
Serious ac	lverse events	(follow-u	o 2 years)									
1	randomised trials	very serious¹	no serious inconsistency	serious <sup>4</sup>	very serious²	none	3/18 (16.7%)	38.9%	RR 0.43 (0.13 to 1.4)	222 fewer per 1000 (from 338 fewer to 156 more)	⊕000 VERY LOW	IMPORTANT

Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias
 Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs
 Absolute effect calculated manually using risk difference as zero events in the studies
 Downgraded by 1 increment as the study does not report withdrawal due to adverse events

MIDs used to assess imprecision were ±66.90
 MIDs used to assess imprecision were ±270.50

# F.1.15 Primary tricuspid regurgitation

2 No studies identified.

# F.136 Secondary heart valve disease – mitral regurgitation or tricuspid regurgitation

4 No studies identified.

5

# F.2 Valve disease with heart failure

# F.271 Primary aortic stenosis

8 Table 45: Clinical evidence profile: ACE-I versus placebo in aortic stenosis

			Quality as	sessment			N	o of patients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	ACE-I	Placebo: primary aortic stenosis	Relative (95% CI)	Absolute		
Exercise 1	xercise tolerance: change in exercise duration (minutes) (follow-up mean 3 days; measured with: semisupine cycle exercise test; Better indicated by lower values)											
	randomised trials	serious <sup>1</sup>	no serious inconsistency	serious <sup>2</sup>	no serious imprecision <sup>3</sup>	none	21	22	-	MD 0 higher (0.31 lower to 0.31 higher)	⊕⊕OO LOW	IMPORTANT
Exercise tolerance: 6-minute walk distance (meters) (follow-up mean 4 weeks; Better indicated by lower values)												
	randomised trials	serious <sup>1</sup>	no serious inconsistency	no serious indirectness	serious <sup>4,5</sup>	none	34	18	-	MD 26 higher (68.89 lower to 120.89 higher)	⊕⊕OO LOW	IMPORTANT

Withdra	Withdrawal due to adverse events (follow-up 2-3 months)													
2	randomised trials		no serious inconsistency	serious <sup>7</sup>	very serious <sup>8</sup>	none	4/59 (6.8%)	2.6%	OR 2.18 (0.34 to 14.17)	29 more per 1000 (from 17 fewer to 248 more)		IMPORTANT		

Heart valve disease: DRAFT FOR CONSULTATION 4 MIDs used to assess imprecision were ±4.0

Clinical evidence profile: ARB versus placebo Table 46:

	Quality assessment  No of Risk of Learning Indicates Learning						_	lo of tients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	ARB	Placebo	Relative (95% CI)	Absolute		
All-cause	All-cause mortality (follow-up 2-12 months)											
1	randomised trials	very serious¹	no serious inconsistency	no serious indirectness	serious <sup>2</sup>	none	0/25 (0%)	3.9%	OR 0.14 (0 to 7.09)	39 fewer per 1000 (from 140 fewer to 63 more) <sup>3</sup>	⊕OOO VERY LOW	CRITICAL
Acute heart failure (follow-up 2-12 months)												
1	randomised trials	very serious¹	no serious inconsistency	serious <sup>4</sup>	very serious <sup>2</sup>	none	1/25 (4%)	0%	OR 7.69 (0.15 to 387.87)	40 more per 1000 (from 63 fewer to 143 more) <sup>3</sup>	⊕OOO VERY LOW	CRITICAL
Exercise t	tolerance: cha	nge from	baseline 6-minute	walking distance	(follow-up 2-12	months; Better in	dicate	ed by low	er values)			
1	randomised trials	serious <sup>5</sup>	no serious inconsistency		no serious imprecision <sup>6</sup>	none	22	21	-	MD 18 lower (48.74 lower to 12.74 higher)	⊕⊕⊕O MODERATE	IMPORTANT
Withdraw	al due to adve	rse events	s (follow-up 2-12 n	nonths)				•				
1	randomised trials	very serious¹	no serious inconsistency	no serious indirectness	very serious <sup>7</sup>	none	2/25 (8%)	7.7%	RR 1.04 (0.16 to 6.83)	3 more per 1000 (from 65 fewer to 449 more)	⊕OOO VERY LOW	IMPORTANT

<sup>&</sup>lt;sup>1</sup> Downgraded by 1 increment because the evidence was at high risk of bias <sup>2</sup> Downgraded by 1 increment because the mean follow-up period was less than 1 month

<sup>&</sup>lt;sup>3</sup> MIDs used to assess imprecision were ±1.0

<sup>&</sup>lt;sup>4</sup> Downgraded by 1 increment because the confidence interval crossed one MID

<sup>&</sup>lt;sup>5</sup> MIDs used to assess imprecision were ±76.0

<sup>&</sup>lt;sup>6</sup> Downgraded by 1 increment because the majority of evidence was at high risk of bias

<sup>7</sup> Downgraded by 1 increment because the mean follow-up period was less than 3 months

<sup>8</sup> Downgraded by 2 increments because the confidence interval crossed both MIDs

15

9

- 10 11 <sup>2</sup> Downgraded by 1 increment because the confidence interval crossed one MID 12 13
  - <sup>4</sup> Absolute effect calculated manually as zero events in one arm of the study

- <sup>1</sup> Downgraded by 2 increments because the evidence was at very high risk of bias <sup>2</sup> Downgraded by 1 increment because the confidence interval crossed one MID
- 3 Absolute effect calculated manually using risk difference as zero events in one arm of the study
- <sup>4</sup> Downgraded by 1 increment because of uncertainty as to the aetiology of reported acute heart failure
- <sup>5</sup> Downgraded by 1 increment because the evidence was at high risk of bias
- <sup>6</sup> MIDs used to assess imprecision were ±74.0
  - <sup>7</sup> Downgraded by 2 increments because the confidence interval crossed both MIDs

#### F.282 **Primary mitral stenosis**

Clinical evidence profile: Beta-blocker versus usual care Table 47:

ubic 7	•		idence prome	. Deta biooi	tor vorsus u	ouur our o						
	Quality assessment  No of Resign Risk of Inconsistency Indirectness Improvious Other									Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Beta- blocker	Usual care	Relative (95% CI)	Absolute		
Hospitalis	sation due to I	heart failu	re (follow-up mear	12 months)								
I	randomised trials		no serious inconsistency	no serious indirectness	serious <sup>2</sup>	none	3/33 (9.1%)	29.4%	RR 0.31 (0.09 to 1.02)	203 fewer per 1000 (from 268 fewer to 6 more)	⊕OOO VERY LOW	CRITICAL
Exercise	tolerance: 6-m	ninute wal	king distance (follo	ow-up 6-12 mont	hs; Better indica	ted by lower value	es)					
1	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision <sup>3</sup>	none	33	34	-	MD 133 higher (121.49 to 144.51 higher)	⊕⊕OO LOW	IMPORTANT
Nithdraw	al due to adve	erse event	s (follow-up mean	12 months)								
1	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	5/44 (11.4%)	0%	OR 8.14 (1.35 to 48.97)	114 more per 1000 (from 13 more to 214 more) <sup>4</sup>	⊕⊕OO LOW	IMPORTANT

<sup>&</sup>lt;sup>1</sup> Downgraded by 2 increments because the evidence was at very high risk of bias

Clinical evidence profile: Beta-blocker versus placebo Table 48:

Quality assessment	No of patients	Effect	Quality	Importance	
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<sup>&</sup>lt;sup>3</sup> MIDs used to assess imprecision were ±15.0

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Beta blocker	Placebo	Relative (95% CI)	Absolute			
Exercise to	Exercise tolerance: treadmill exercise time (minutes) to exhaustion (follow-up 1-4 weeks; Better indicated by lower values)												
3		very serious <sup>1</sup>	serious <sup>2</sup>	serious³	very serious <sup>4,5</sup>	none	42	42	-	MD 0.33 higher (1.09 lower to 1.75higher)	⊕OOO VERY LOW	IMPORTANT	
Exercise to	Exercise tolerance: Pulmonary capillary wedge pressure after exercise (follow-up mean 6 months; Better indicated by lower values)												
1	randomised trials		no serious inconsistency		no serious imprecision <sup>8</sup>	none	13	13	-	MD 14.8 lower (21.71 to 7.89 lower)	⊕⊕OO LOW	IMPORTANT	

Clinical evidence profile: Beta-blocker versus calcium channel blocker Table 49:

Quality assessment							No of patients		Effect		Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Beta blocker	Calcium channel blocker	Relative (95% CI)	Absolute		
Exercise tolerance: total effort time on treadmill exercise test (follow-up mean 3 months; Better indicated by lower values)												
1		- ,	no serious inconsistency	no serious indirectness	serious <sup>2,3</sup>	none	40	40	-	MD 50 lower (97.99 to 2.01 lower) <sup>4</sup>	⊕OOO VERY LOW	IMPORTANT
Withdrawal due to adverse events (follow-up mean 3 months)												
1	trials			indirectness		none	0/40 (0%)	0%	-RD 0 (-0.048 to 0.048)	0 fewer per 1000 (from 48 fewer to 48 more) <sup>7</sup>	⊕⊕OO LOW	IMPORTANT

Downgraded by 2 increments because the evidence was at very high risk of bias

10

<sup>&</sup>lt;sup>1</sup> Downgraded by 2 increments because the majority of the evidence was at very high risk of bias <sup>2</sup> Downgraded by one increment because the I2 = 74% and heterogeneity was not explained by subgroup analyses.

<sup>&</sup>lt;sup>3</sup> Downgraded by 1 increment because the mean follow-up period is less than 1 month

<sup>&</sup>lt;sup>4</sup> Downgraded by 2 increments because the confidence interval crossed both MIDs

<sup>&</sup>lt;sup>5</sup> MIDs used to assess imprecision were ±0.9

<sup>&</sup>lt;sup>6</sup> Downgraded by 1 increment because the evidence was at high risk of bias

<sup>&</sup>lt;sup>7</sup> Downgraded by 1 increment because the outcome is a surrogate measure

<sup>&</sup>lt;sup>8</sup> MIDs used to assess imprecision were ±5.35

<sup>&</sup>lt;sup>2</sup> Downgraded by 1 increment because the confidence interval crossed one MID

<sup>&</sup>lt;sup>3</sup> MIDs used to assess imprecision were ±60.0

<sup>&</sup>lt;sup>4</sup> Baseline total effort time not matched

<sup>&</sup>lt;sup>5</sup> Downgraded by 1 increment because the evidence was at high risk of bias

<sup>&</sup>lt;sup>6</sup> Downgraded by 1 increments because sample size was >70 and <350 (imprecision was assessed based on sample size as zero events in both arms of the study)

<sup>&</sup>lt;sup>7</sup> Absolute effect calculated manually as zero events in both arms of the study

#### F.2.13 Secondary heart valve disease (mitral regurgitation and tricuspid regurgitation)

Table 50: Clinical evidence profile: ACE-I versus placebo in secondary heart valve disease

					<u>-</u>			,,	L			
Quality assessment								No of patients		Effect		Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	ACE-	Placebo: Secondary heart valve disease – mitral regurgitation or tricuspid regurgitation	Relative (95% CI)	Absolute	Quality	Importance
Cardiac r	Cardiac mortality (follow-up mean 12 weeks)											
	randomised trials		no serious inconsistency	no serious indirectness	serious <sup>2</sup>	none	0/14 (0%)	7.1%	OR 0.14 (0 to 6.82)	71 fewer per 1000 (from 248 fewer to 106 more) <sup>3</sup>		CRITICAL
Quality o	Quality of life: Duke activity index score (follow-up mean 12 weeks; range of scores: 2.75-58.2; Better indicated by lower values)											
		- ,	no serious inconsistency	serious <sup>5</sup>	serious <sup>2,6</sup>	none	10	13	-	MD 6.7 higher (0.97 lower to 14.37 higher)	⊕OOO VERY LOW	CRITICAL
Withdrawal due to adverse events (follow-up mean 3 months)												
1	randomised trials		no serious inconsistency	no serious indirectness	very serious <sup>7</sup>	none	0/14 (0%)	0%	-RD 0 (- 0.133 to 0.133)	0 fewer per 1000 (from 133 fewer to 133 more) <sup>8</sup>	⊕OOO VERY LOW	IMPORTANT

<sup>&</sup>lt;sup>1</sup> Downgraded by 1 increment because the evidence was at high risk of bias
<sup>2</sup> Downgraded by 1 increment because the confidence interval crossed one MID
<sup>3</sup> Absolute effect calculated manually using risk difference as zero events in one arm of the study

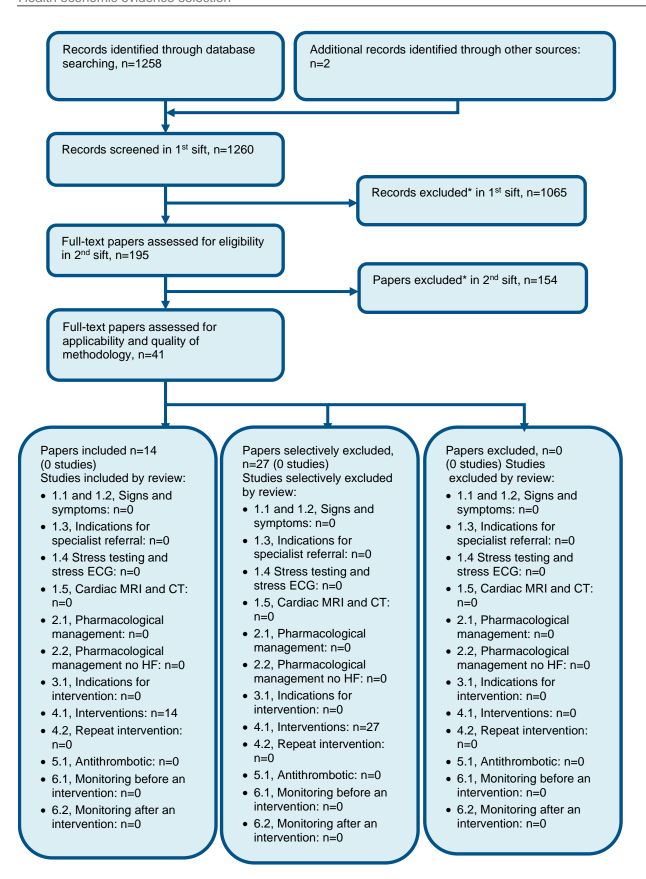
<sup>&</sup>lt;sup>4</sup> Downgraded by 2 increments because the evidence was at very high risk of bias

<sup>&</sup>lt;sup>5</sup> Downgraded by 1 increment because the reported measure only reports physical activity rather than other aspects of quality of life <sup>6</sup> MIDs used to assess imprecision were ±4.7

<sup>&</sup>lt;sup>7</sup> Downgraded by 2 increments because sample size was <70 (imprecision was assessed based on sample size as zero events in both arms of the study) <sup>8</sup> Absolute effect calculated manually as zero events in both arms of the study

# Appendix G: Health economic evidence

<sub>2</sub> selection



<sup>\*</sup> Non-relevant population, intervention, comparison, design or setting; non-English language

# **Appendix H: Health economic evidence tables**

H.4 Valve disease without heart failure

4 None.

H.2 Valve disease with heart failure

6 None.

7

# 2 Appendix I: Excluded studies

# I.4 Valve disease without heart failure

### I.15 Excluded clinical studies

1

3

### 6 Table 51: Studies excluded from the clinical review

Study	Exclusion reason						
Agnihotri 2017 <sup>1</sup>	Could not be retrieved						
Agrawal 2016 <sup>2</sup>	Incorrect interventions. Uses Ivabradine that is not included in ou scope. Included patients with mild valve disease						
Ahmed 2002 <sup>3</sup>	Incorrect study design. Mixed population <75% HVD						
Ahuja 1989 <sup>5</sup>	Incorrect study design						
Alan 2002 <sup>6</sup>	Not review population						
Andersson 2002 <sup>7</sup>	Incorrect study design. Not guideline condition. Not review population. Mixed population <75% HVD						
Andersson 2017 <sup>8</sup>	Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear. Inappropriate comparison. References checked and extracted						
Andrus 2007 <sup>9</sup>	Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear. Systematic review: literature search not sufficiently rigorous. Systematic review: study designs inappropriate. Not review population. Inappropriate comparison. Incorrect interventions						
Antonini-Canterin 2006 <sup>10</sup>	Protocol only						
Aumont 1990 <sup>11</sup>	Not in English language						
Bassan 1987 <sup>15</sup>	Not review population						
Bechler-lisińska 1990 <sup>16</sup>	Not in English language						
Bergstrom 2004 <sup>17</sup>	Not guideline condition. Not review population						
Borer 1978 <sup>18</sup>	Incorrect study design. Inappropriate comparison. Incorrect interventions						
Bornheimer 1982 <sup>19</sup>	Incorrect study design. Inappropriate comparison. No control group. Severity not mentioned						
Butrous 1986 <sup>22</sup>	Incorrect study design. Not review population.						
Capucci 1981 <sup>23</sup>	Not in English language						
Carabello 2010 <sup>24</sup>	Correspondence only						
Chockalingam 2004P <sup>29</sup>	Not review population						
Choi 2015 <sup>30</sup>	Systematic review: study designs inappropriate. Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear						
Chua 2006 <sup>31</sup>	Systematic review: study designs inappropriate. Systematic review: quality assessment is inadequate						
Cicoira 2002 <sup>32</sup>	Not guideline condition. Not review population						
Clark 1983 <sup>33</sup>	Not guideline condition						

Cleland 2006 <sup>34</sup>	Not guideline condition. Not review population	
Cohen 1968 <sup>35</sup>	Not guideline condition. Not review population. Inappropriate	
Crawford 1989 <sup>37</sup>	Incorrect study design. Incorrect interventions	
Dalsgaard 2014 <sup>38</sup>	Not review population	
De Vicchis 2013 <sup>39</sup>	Systematic review; quality assessment is inadequate	
Demirbag 2003 <sup>40</sup>	Not in English language	
De vecchis 2013 <sup>39</sup>	Systematic review: quality assessment is inadequate. Systematic review: study designs inappropriate	
Eichhorn 2001 <sup>42</sup>	Not guideline condition. Not review population	
Eleid 2013 <sup>43</sup>	Incorrect study design. Inappropriate comparison. Incorrect interventions	
Ennis 2010 <sup>44</sup>	No appropriate outcomes reported	
Erbel 1978 <sup>45</sup>	Not in English language	
Erbel 1979 <sup>46</sup>	No appropriate outcomes reported	
Ergun 2016 <sup>47</sup>	Incorrect study design. Letter	
Eskandr 2018 <sup>48</sup>	Not review population. Medical management in the intra- or post- operative period. Incorrect line of therapy. Incorrect interventions	
Ge 2010 <sup>50</sup>	Systematic review: study designs inappropriate. Systematic review: quality assessment is inadequate	
Ghadimi 2019 <sup>51</sup>	Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Incorrect line of therapy	
Ghiringhelli 1990 <sup>52</sup>	Incorrect study design. Incorrect interventions	
Giles 1988 <sup>53</sup>	Not guideline condition. Not review population. Mixed population <75% HVD	
Giunta 1993 <sup>54</sup>	Not guideline condition. Not review population	
Gottlieb 2018 <sup>55</sup>	Systematic review is not relevant to review question or unclear PICO. Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear. Inappropriate comparison. Incorrect line of therapy	
Grayburn 2000 <sup>56</sup>	Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear. Inappropriate comparison. Incorrect line of therapy	
Greenberg 1994 <sup>57</sup>	Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear. Incorrect line of therapy	
Greve 2008 <sup>61</sup>	Systematic review: study designs inappropriate. Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear	
Gupta 2001 <sup>62</sup>	Incorrect interventions	
Hachenberg 1997 <sup>63</sup>	Incorrect interventions	
Hamilton-craig 2009 <sup>64</sup>	Correspondence only	
Han 2018 <sup>65</sup>	Systematic review: methods are not adequate/unclear. Systematic review is not relevant to review question or unclear PICO. Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Inappropriate comparison. Incorrect line of therapy. Incorrect interventions	

Helske-Suihko 2015 <sup>67</sup>	Not review population
Henriquez 2009 <sup>68</sup>	Systematic review is not relevant to review question or unclear PICO. Incorrect interventions
Hjalmarson 1991 <sup>69</sup>	Not guideline condition. Not review population. Systematic review is not relevant to review question or unclear PICO. Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear. Inappropriate comparison
Hjalmarson 1994 <sup>70</sup>	Systematic review is not relevant to review question or unclear PICO. Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear. Not guideline condition. Not review population. Inappropriate comparison
Hongning 2014 <sup>72</sup>	No appropriate outcomes reported
Host 1997 <sup>73</sup>	Incorrect outcomes
Hung 2002 <sup>75</sup>	Incorrect study design. Not guideline condition. Not review population
Ibragimova 2018 <sup>76</sup>	Not available in English language
Inano 2007 <sup>77</sup>	Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear
Jirasirirojanakorn 1998 <sup>78</sup>	Incorrect population: combines different types of heart valve disease
Kang 2019 <sup>80</sup>	Incorrect interventions
Kasama 2007 <sup>81</sup>	Mixed population <75% HVD
Kelbaek 199682	Not review population
Keren 1992 <sup>83</sup>	Not review population
Keren 1994 <sup>84</sup>	Not guideline condition. Not review population
Kesaniemi 200785	Commentary only
Kleaveland 198686	Not review population. Mixed population <75% heart failure. Incorrect interventions
Klein 1985 <sup>87</sup>	Not review population
Klugmann 1983 <sup>88</sup>	Not guideline condition. Not review population. Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear. Inappropriate comparison
Korewicki 199089	Not available in English language
Kumar 1994 <sup>90</sup>	Not review population
Lanas 1995 <sup>91</sup>	Not available in English language
Lanas 1996 <sup>93</sup>	Not available in English language
Lanas 1998 <sup>92</sup>	Not available in English language
Leenen 199194	Incorrect study design
Legault 1996 <sup>95</sup>	Not review population. Medical management in the intra- or post- operative period
Levine 1998 <sup>96</sup>	Not review population. Incorrect study design. Inappropriate comparison
Lin 1994 <sup>98</sup>	Not guideline condition. Not review population. Incorrect interventions
Lin 1994 <sup>99</sup>	Not review population. Incorrect interventions
Lin 2011 <sup>97</sup>	Not review population. Systematic review is not relevant to review question or unclear PICO. Systematic review: literature search not

	sufficiently rigorous. Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear
Littler 1995 <sup>100</sup>	Not guideline condition. Not review population
Loomba 2010 <sup>101</sup>	Systematic review: study designs inappropriate. Systematic review: methods are not adequate/unclear. Systematic review: quality assessment is inadequate
Lowes 1999 <sup>102</sup>	Not guideline condition. Not review population
Mahajerin 2007 <sup>103</sup>	Not review population
Mardikar 1995 <sup>105</sup>	Incorrect study design
Memon 2016 <sup>106</sup>	Systematic review: quality assessment is inadequate. Systematic review: study designs inappropriate
Misra 1989 <sup>107</sup>	Incorrect study design. Severity of heart disease not mentioned
Mizuno 2002 <sup>108</sup>	Not guideline condition. Not review population
Moura 2007 <sup>109</sup>	Incorrect study design
Muhammad 2016 <sup>110</sup>	Incorrect interventions. Mild-to-moderate heart valve disease
Nagatomo 2007 <sup>111</sup>	Not review population. Not guideline condition
Nikitin 1998 <sup>113</sup>	Not guideline condition. Not review population. Inappropriate comparison
Novo 2011 <sup>114</sup>	Systematic review: study designs inappropriate. Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear
Nyolczas 2017 <sup>115</sup>	Not review population. Systematic review is not relevant to review question or unclear PICO. Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear. Inappropriate comparison
Olsen 2004 <sup>116</sup>	Not review population
Olsson 2009 <sup>117</sup>	Systematic review: literature search not sufficiently rigorous.  Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear
Packer 1983 <sup>118</sup>	Not guideline condition. Not review population. Systematic review: methods are not adequate/unclear. Systematic review: quality assessment is inadequate. Systematic review: literature search not sufficiently rigorous. Systematic review is not relevant to review question or unclear PICO. Inappropriate comparison
Panahi 2013 <sup>119</sup>	Not review population
Parakh 2012 <sup>120</sup>	Not review population. Incorrect interventions. Mild to moderate mitral stenosis
Park 2016 <sup>121</sup>	Medical management in the intra- or post-operative period
Parolari 2011 <sup>122</sup>	Systematic review: study designs inappropriate. Systematic review: quality assessment is inadequate
Patel 1995 <sup>123</sup>	Not review population
Pedersen 2008 <sup>124</sup>	Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear. Systematic review: study designs inappropriate. Systematic review: literature search not sufficiently rigorous
Quinn 2005 <sup>125</sup>	Correspondence only
Rajesh 2016 <sup>126</sup>	Incorrect interventions
Ramos 2018 <sup>127</sup>	Incorrect interventions. Systematic review: methods are not adequate/unclear. Systematic review: quality assessment is inadequate. Systematic review: literature search not sufficiently
	rigorous

Rivera 2003 <sup>128</sup>	Not review population. Mixed population <75% heart failure. Mild mitral regurgitation
Rosenhek 2008 <sup>130</sup>	Systematic review: study designs inappropriate. Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear
Roth 1993 <sup>133</sup>	Medical management in the intra- or post-operative period. Incorrect interventions
Rothlisberger 1993 <sup>134</sup>	Not review population
Rothlisberger 1994 <sup>135</sup>	Not review population
Ruiz Ros 1999 <sup>136</sup>	Medical management in the intra- or post-operative period. Not review population
Saeed 2020 <sup>137</sup>	Incorrect study design
Saggu 2015 <sup>138</sup>	Incorrect interventions. Mild-to-moderate mitral stenosis
Sahebkar 2012 <sup>139</sup>	Conference abstract only
Sahoo 2016 <sup>140</sup>	No protocol outcomes
Salas 2012 <sup>141</sup>	Not review population. Systematic review is not relevant to review question or unclear PICO. Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear. Inappropriate comparison
Saltissi 1983 <sup>142</sup>	Not review population. Mitral valve prolapse without regurgitation
Sanada 2007 <sup>144</sup>	Not guideline condition. Not review population. Incorrect interventions
Seneviratne 1994 <sup>147</sup>	Not review population
Shah 2012 <sup>148</sup>	Not review population. Systematic review is not relevant to review question or unclear PICO. Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear
Shen 1995 <sup>149</sup>	Not available in English language
Shu 2005 <sup>150</sup>	Not review population
Slipczuk 2016 <sup>151</sup>	Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear
Sondergaard 2000 <sup>152</sup>	Not review population
Stewart 2008 <sup>155</sup>	No appropriate outcomes reported
Stewart 2009 <sup>153</sup>	Systematic review is not relevant to review question or unclear PICO. Systematic review: study designs inappropriate. Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear
Stoll 1995 <sup>156</sup>	Mild heart valve disease
Strauss 2012 <sup>157</sup>	Not review population. Systematic review is not relevant to review question or unclear PICO. Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear
Takagi 2009 <sup>159</sup>	Letter only
Takagi 2019 <sup>158</sup>	Not review population
Takahama 2018 <sup>160</sup>	Not guideline condition. Not review population. Inappropriate comparison
Tan 1998 <sup>161</sup>	Not available in English language
Tendera 1987 <sup>162</sup>	Not available in English language

Teo 2011 <sup>163</sup>	Systematic review: quality assessment is inadequate
Thilly 2003 <sup>164</sup>	Not guideline condition. Not review population. Inappropriate comparison
Tjon 1990 <sup>165</sup>	Not available in English language
Tourmousoglou 2008 <sup>166</sup>	Systematic review: methods are not adequate/unclear. Systematic review: quality assessment is inadequate. Systematic review: literature search not sufficiently rigorous. Systematic review: study designs inappropriate
Tschirkov 1992 <sup>167</sup>	Medical management in the intra- or post-operative period
Van der Linde 2011 <sup>168</sup>	Not review population
Venegas 1992 <sup>169</sup>	Not available in English language
Venegas 1992 <sup>170</sup>	Not available in English language
Vizzardi 2010 <sup>171</sup>	Not guideline condition. Not review population
Vonder Muhll 2004 <sup>172</sup>	Not review population. Systematic review is not relevant to review question or unclear PICO. Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear
Waagstein 2003 <sup>173</sup>	Not guideline condition. Not review population
Wagner 2003 <sup>174</sup>	Medical management in the intra- or post-operative period. Incorrect study design. Incorrect interventions
Wenaweser 2011 <sup>175</sup>	Incorrect study design. Incorrect interventions
Wisenbaugh 1991 <sup>176</sup>	Incorrect study design. Incorrect interventions
Witczak 2008 <sup>178</sup>	Medical management in the intra- or post-operative period. Mixed population <75% HVD
Yurpolskaya 1996 <sup>179</sup>	Not available in English language
Zhao 2016 <sup>180</sup>	Systematic review: study designs inappropriate. Systematic review: quality assessment is inadequate
Zhao 2016 <sup>181</sup>	Systematic review: quality assessment is inadequate. Systematic review: study designs inappropriate
Zhou 2008 <sup>182</sup>	Not available in English language

### I.132 Excluded health economic studies

- 4 Published health economic studies that met the inclusion criteria (relevant population,
- 5 comparators, economic study design, published 2004 or later and not from non-OECD
  - country or USA) but that were excluded following appraisal of applicability and
- 7 methodological quality are listed below. See the health economic protocol for more details.
- 8 None.

9

# I.2 Valve disease with heart failure

# I.23 Excluded clinical studies

#### 4 Table 52: Studies excluded from the clinical review

Study	Exclusion reason
Agnihotri 2017 <sup>1</sup>	Could not be retrieved
Agrawal 2016 <sup>2</sup>	Incorrect interventions. Uses Ivabradine that is not included in our scope. Included patients with mild valve disease
Ahmed 2002 <sup>3</sup>	Incorrect study design. Mixed population <75% HVD
Ahmed 2012 <sup>4</sup>	Mixed population <75% heart failure
Ahuja 1989 <sup>5</sup>	Incorrect study design
Andersson 2002 <sup>7</sup>	Incorrect study design. Not guideline condition. Not review population. Mixed population <75% HVD
Andersson 2017 <sup>8</sup>	Systematic review: literature search not sufficiently rigorous.  Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear. Inappropriate comparison. References checked and extracted
Andrus 2007 <sup>9</sup>	Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear. Systematic review: literature search not sufficiently rigorous. Systematic review: study designs inappropriate. Not review population. Inappropriate comparison. Incorrect interventions
Aumont 1990 <sup>11</sup>	Not in English
Balmforth 2019 <sup>12</sup>	Not review population. Incorrect interventions.
Banaszewski 1998 <sup>13</sup>	Incorrect study design. Not review population. Mixed population <75% heart failure
Bechler-Lisińska 1990 <sup>16</sup>	Not in English language
Bergstrom 2004 <sup>17</sup>	Not guideline condition. Not review population
Borer 1978 <sup>18</sup>	Incorrect study design. Inappropriate comparison. Incorrect interventions
Bornheimer 1982 <sup>19</sup>	Incorrect study design. Inappropriate comparison. No control group. Severity not mentioned
Broch 2016 <sup>20</sup>	Not review population. No patients with heart failure
Bull 2015 <sup>21</sup>	Not review population. No participants with heart failure
Butrous 1986 <sup>22</sup>	Incorrect study design. Not review population. No participants with heart failure. No mention of severity
Capucci 1981 <sup>23</sup>	Not in English language
Cicoira 2002 <sup>32</sup>	Not guideline condition. Not review population
Clark 1983 <sup>33</sup>	Not guideline condition
Cleland 2006 <sup>34</sup>	Not guideline condition. Not review population
Cohen 1968 <sup>35</sup>	Not guideline condition. Not review population. Inappropriate comparison
Crawford 1989 <sup>37</sup>	Incorrect study design. Incorrect interventions

Eichhorn 2001 <sup>42</sup>	Not guideline condition. Not review population
Eleid 2013 <sup>43</sup>	Incorrect study design. Inappropriate comparison. Incorrect interventions
Ennis 2010 <sup>44</sup>	Mixed population <75% heart failure
Erbel 1978 <sup>45</sup>	Not available in English language
Erbel 1979 <sup>46</sup>	Not review population. No mention of severity
Ergun 2016 <sup>47</sup>	Incorrect study design. Letter
Eskandr 2018 <sup>48</sup>	Not review population. Medical management in the intra- or post- operative period. Incorrect line of therapy. Incorrect interventions
Evangelista 2005 <sup>49</sup>	Not review population
Ghadimi 2019 <sup>51</sup>	Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Incorrect line of therapy
Ghiringhelli 1990 <sup>52</sup>	Incorrect study design. Incorrect interventions
Giles 1988 <sup>53</sup>	Not guideline condition. Not review population. Mixed population <75% HVD
Giunta 1993 <sup>54</sup>	Not guideline condition. Not review population
Gottlieb 2018 <sup>55</sup>	Systematic review is not relevant to review question or unclear PICO. Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear. Inappropriate comparison. Incorrect line of therapy
Grayburn 2000 <sup>56</sup>	Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear. Inappropriate comparison. Incorrect line of therapy
Greenberg 1994 <sup>57</sup>	Systematic review: literature search not sufficiently rigorous.  Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear. Incorrect line of therapy
Gupta 2001 <sup>62</sup>	Incorrect interventions
Hachenberg 1997 <sup>63</sup>	Incorrect interventions
Han 2018 <sup>65</sup>	Systematic review: methods are not adequate/unclear. Systematic review is not relevant to review question or unclear PICO. Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Inappropriate comparison. Incorrect line of therapy. Incorrect interventions
Hansson 2017 <sup>66</sup>	Not review population. Incorrect line of therapy
Henriquez 2009 <sup>68</sup>	Systematic review is not relevant to review question or unclear PICO. Incorrect interventions
Hjalmarson 1991 <sup>69</sup>	Not guideline condition. Not review population. Systematic review is not relevant to review question or unclear PICO. Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear. Inappropriate comparison
Hjalmarson 1994 <sup>70</sup>	Systematic review is not relevant to review question or unclear PICO. Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear. Not guideline condition. Not review population. Inappropriate comparison
Hongning 2014 <sup>72</sup>	Mixed population <75% heart failure

Host 1997 <sup>73</sup>	Incorrect outcomes
Hung 2002 <sup>75</sup>	Incorrect study design. Not guideline condition. Not review
Hully 2002.	population
Ibragimova 2018 <sup>76</sup>	Not available in English language
Inano 2007 <sup>77</sup>	Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear
Jirasirirojanakorn 1998 <sup>78</sup>	Incorrect population: combines different types of heart valve disease
Kang 201980	Incorrect interventions
Kasama 200781	Mixed population <75% HVD
Kelbaek 199682	Incorrect outcomes
Keren 199283	Majority of participants had mild valve disease
Keren 199484	Not guideline condition. Not review population
Kleaveland 198686	Not review population. Mixed population <75% heart failure. Incorrect interventions
Klugmann 198388	Not guideline condition. Not review population. Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear. Inappropriate comparison
Korewicki 199089	Not available in English language
Lanas 199591	Not available in English language
Lanas 1996 <sup>93</sup>	Not available in English language
Lanas 1998 <sup>92</sup>	Not available in English language
Legault 1996 <sup>95</sup>	Not review population. Medical management in the intra- or post- operative period
Levine 1998 <sup>96</sup>	Not review population. Incorrect study design. Inappropriate comparison
Lin 1994 <sup>98</sup>	Not guideline condition. Not review population. Incorrect interventions
Lin 1994 <sup>99</sup>	Not review population. Incorrect interventions
Lin 2011 <sup>97</sup>	Not review population. Systematic review is not relevant to review question or unclear PICO. Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear
Littler 1995 <sup>100</sup>	Not guideline condition. Not review population
Lowes 1999 <sup>102</sup>	Not guideline condition. Not review population
Mahajerin 2007 <sup>103</sup>	Not review population
Marcotte 1997 <sup>104</sup>	Not review population
Mardikar 1995 <sup>105</sup>	Incorrect study design
Misra 1989 <sup>107</sup>	Incorrect study design. Severity of heart disease not mentioned
Mizuno 2002 <sup>108</sup>	Not guideline condition. Not review population
Muhammad 2016 <sup>110</sup>	Incorrect interventions. Mild-to-moderate heart valve disease
Nagatomo 2007 <sup>111</sup>	Not review population. Not guideline condition
Nikitin 1998 <sup>113</sup>	Not guideline condition. Not review population. Inappropriate comparison

Nyolczas 2017 <sup>115</sup>	Not review population. Systematic review is not relevant to review question or unclear PICO. Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear. Inappropriate comparison
Olsen 2004 <sup>116</sup>	Not review population
Packer 1983 <sup>118</sup>	Not guideline condition. Not review population. Systematic review: methods are not adequate/unclear. Systematic review: quality assessment is inadequate. Systematic review: literature search not sufficiently rigorous. Systematic review is not relevant to review question or unclear PICO. Inappropriate comparison
Parakh 2012 <sup>120</sup>	Not review population. Incorrect interventions. Mild to moderate mitral stenosis
Rajesh 2016 <sup>126</sup>	Incorrect interventions
Ramos 2018 <sup>127</sup>	Incorrect interventions. Systematic review: methods are not adequate/unclear. Systematic review: quality assessment is inadequate. Systematic review: literature search not sufficiently rigorous
Rivera 2003 <sup>128</sup>	Not review population. Mixed population <75% heart failure. Mild mitral regurgitation
Roberts 2018 <sup>129</sup>	Not review population
Roth 1993 <sup>133</sup>	Medical management in the intra- or post-operative period. Incorrect interventions
Rothlisberger 1993 <sup>134</sup>	Not review population
Rothlisberger 1994 <sup>135</sup>	Not review population
Ruiz Ros 1999 <sup>136</sup>	Medical management in the intra- or post-operative period. Not review population
Saeed 2020 <sup>137</sup>	Incorrect study design
Saggu 2015 <sup>138</sup>	Incorrect interventions. Mild-to-moderate mitral stenosis
Sahoo 2016 <sup>140</sup>	No protocol outcomes
Salas 2012 <sup>141</sup>	Not review population. Systematic review is not relevant to review question or unclear PICO. Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear. Inappropriate comparison
Saltissi 1983 <sup>142</sup>	Not review population. Mitral valve prolapse without regurgitation
Sampaio 2005 <sup>143</sup>	Mixed population <75% heart failure
Sanada 2007 <sup>144</sup>	Not guideline condition. Not review population. Incorrect interventions
Scognamiglio 1990 <sup>145</sup>	Not review population
Scognamiglio 1994 <sup>146</sup>	Not review population
Shah 2012 <sup>148</sup>	Not review population. Systematic review is not relevant to review question or unclear PICO. Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear
Shen 1995 <sup>149</sup>	Not available in English language
Slipczuk 2016 <sup>151</sup>	Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear
Sondergaard 2000 <sup>152</sup>	Not review population

Stewart 2008 <sup>154</sup>	Not review population
Stewart 2008 <sup>155</sup>	Mixed population <75% heart failure
Stoll 1995 <sup>156</sup>	Mild heart valve disease
Strauss 2012 <sup>157</sup>	Not review population. Systematic review is not relevant to review question or unclear PICO. Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear
Takahama 2018 <sup>160</sup>	Not guideline condition. Not review population. Inappropriate comparison
Tan 1998 <sup>161</sup>	Not available in English language
Tendera 1987 <sup>162</sup>	Not available in English language
Thilly 2003 <sup>164</sup>	Not guideline condition. Not review population. Inappropriate comparison
Tjon 1990 <sup>165</sup>	Not available in English language
Tschirkov 1992 <sup>167</sup>	Medical management in the intra- or post-operative period
Venegas 1992 <sup>169</sup>	Not available in English language
Venegas 1992 <sup>170</sup>	Not available in English language
Vizzardi 2010 <sup>171</sup>	Not guideline condition. Not review population
Vonder Muhll 2004 <sup>172</sup>	Not review population. Systematic review is not relevant to review question or unclear PICO. Systematic review: literature search not sufficiently rigorous. Systematic review: quality assessment is inadequate. Systematic review: methods are not adequate/unclear
Waagstein 2003 <sup>173</sup>	Not guideline condition. Not review population
Wagner 2003 <sup>174</sup>	Medical management in the intra- or post-operative period. Incorrect study design. Incorrect interventions
Wenaweser 2011 <sup>175</sup>	Incorrect study design. Incorrect interventions
Wisenbaugh 1991 <sup>176</sup>	Incorrect study design. Incorrect interventions
Wisenbaugh 1994 <sup>177</sup>	Mixed population <75% heart failure. Mean NYHA class 1.3
Witczak 2008 <sup>178</sup>	Medical management in the intra- or post-operative period. Mixed population <75% HVD
Yurpolskaya 1996 <sup>179</sup>	Not available in English language
Zhou 2008 <sup>182</sup>	Not available in English language

#### I.22 Excluded health economic studies

- 3 Published health economic studies that met the inclusion criteria (relevant population,
- 4 comparators, economic study design, published 2004 or later and not from non-OECD
- 5 country or USA) but that were excluded following appraisal of applicability and
- 6 methodological quality are listed below. See the health economic protocol for more details.
- 7 None.

# Appendix J: Research recommendations

# J.1 Heart valve disease without concomitant heart failure

#### J.13 Research recommendation

- **J.14** What is the clinical and cost-effectiveness of ACE inhibitors, angiotensin-II receptor
  - 5 antagonists, beta-blockers and calcium channel blockers, including compared with placebo,
  - 6 for adults with aortic regurgitation?

## J.172 Why this is important

13

15

- 8 There are two aspects to this:
- The tolerability and secondary effect on aortic regurgitation of these drugs when taken for other purposes as for example for systemic hypertension or for angina or arrhythmia
   The perceived by some potential role of these drugs in delaying progression of aortic regurgitation of these drugs when taken for other purposes as for example for systemic hypertension or for angina or arrhythmia
  - 2. The perceived by some potential role of these drugs in delaying progression of aortic regurgitation or the consequences of it on symptoms and on the left ventricle.

#### J.143 Rationale for research recommendation

Importance to 'patients' or the population	<ol> <li>Adjust medication appropriately when aortic regurgitation coexists</li> <li>Offer medication appropriately of demonstrated to have a benefit or contraindicate if harm.</li> </ol>
Relevance to NICE guidance	There was very limited evidence on ACE-Is, ARBs, beta-blockers and calcium channel blockers for aortic regurgitation. Additional evidence would enable recommendations to be made.
Relevance to the NHS	Allow standardisation of care and reduce cost by delaying need for intervention if this is proved to be the case.  This is a key area of concern in current UK practice as there is uncertainty about whether pharmacological management is required for people with heart valve disease to prevent progression or delay consequences of the disease and the effect of pharmacological treatment given for other conditions in those that also have heart valve disease. More specifically, there is uncertainty as to whether medications for the management of systemic hypertensions are more poorly tolerated in the presence of valve disease. In addition, there is uncertainty as to whether medications will delay the consequences of valve disease, for example symptoms.
National priorities	None known

Current evidence base	There was insufficient evidence to draw conclusions about the relative benefits and harms of ACE-I, ARB, beta-blockers and calcium channel blockers based on the evidence available. The evidence was based on a small number of studies with a small number of participants. A lot of the studies were historical and so may not reflect current practice.
Equality considerations	Standardisation of care

# J.124 Modified PICO table

Population	Inclusion Adults aged 18 years and over with diagnosed at least moderate aortic regurgitation and no current indication for intervention  Exclusion Pharmacological management in children (17 years and under)
	People with congenital heart valve disease, except bicuspid aortic valve disease.  Known contraindication for the assessed drug
Intervention	<ul> <li>Angiotensin-converting enzyme (ACE) inhibitors</li> <li>Angiotensin-II receptor blockers (ARBs)</li> <li>Beta-blockers</li> <li>Calcium channel blockers</li> <li>Any combination of 2 or more of the above</li> </ul>
Comparator	<ul> <li>Placebo or no treatment (usual care)</li> <li>Other active comparator listed above, including combinations</li> </ul>
Outcome	All-cause mortality at ≥12 months; Cardiac mortality at ≥12 months; Health-related quality of life at 6 months and ≥12 months; Onset of symptoms or progression in NYHA class at ≥12 months; Evidence of HVD progression on imaging (worsening of disease severity) at ≥ 12 months and Need for heart valve intervention (surgical or transcatheter) at ≥12 months  Secondary outcomes:  Exercise tolerance and Withdrawal from the trial due to adverse events at 6 and 12 months
Study design Timeframe	Randomised controlled trial Long term
Timenanie	Long term

#### J.1.5 Research recommendation

**J.126** What is the clinical and cost-effectiveness of ACE inhibitors, beta-blockers and diuretics for adults with severe aortic stenosis?

## J.147 Why this is important

- 5 To assess the tolerability and secondary effects on patients with severe aortic stenosis of
- 6 these drugs when taken for other purposes as for example for systemic hypertension,
- 7 angina, arrhythmia or heart failure symptoms and signs. To assess the potential role of ACE
- 8 inhibitors in delaying progression of consequences of aortic stenosis on the left ventricle,
- 9 consequences that lead to heart failure with preserved ejection fraction, particularly in case
- 10 of coexistence of aortic stenosis and systemic hypertension.

#### J.1.8 Rationale for research recommendation

Importance to 'patients' or the population	Robust evidence covering this area would allow improved management of coexistent conditions in those with aortic stenosis.
Relevance to NICE guidance	Only very limited evidence was available on ACE inhibitors, beta-blockers and diuretics, with studies having small numbers of participants and low event rates across the study periods. It was therefore not possible to draw conclusions about the relative benefits and harms. Additional evidence may enable recommendations to be made in the future.
Relevance to the NHS	ACE inhibitors, beta-blockers and diuretics are widely used in clinical practice for several conditions that can coexist with aortic stenosis. Robust evidence in this population may inform future recommendations about whether these drugs are appropriate in those that have severe aortic stenosis.
	This is a key area of concern in current UK practice as there is uncertainty about whether pharmacological management is required for people with heart valve disease to prevent progression or delay consequences of the disease and the effect of pharmacological treatment given for other conditions in those that also have heart valve disease. More specifically, there is uncertainty as to whether medications for the management of systemic hypertensions are more poorly tolerated in the presence of valve disease. In addition, there is uncertainty as to whether medications will delay the consequences of valve disease, for example symptoms.
National priorities	None known
Current evidence base	There was insufficient evidence to draw conclusions about the relative benefits and

	harms of ACE-I, beta-blockers and diuretics based on the evidence available. While ACE-I and beta-blockers showed clinically important harms, the evidence was collected from small studies that did not show a large enough difference in effect for recommendations to be made.
Equality considerations	Patients with severe aortic stenosis represent a subgroup of patients with systemic hypertension, angina, arrhythmia or heart failure that may necessitate adjustment of the treatment offered to the rest of the group.

### J.129 Modified PICO table

3

Population	Adults aged 18 years and over with diagnosed with severe aortic stenosis, including bicuspid
Intervention	<ul> <li>Angiotensin-converting enzyme (ACE) inhibitors</li> <li>Beta blockers</li> <li>Diuretics</li> <li>Any combination of 2 or more of the above</li> </ul>
Comparator	<ul> <li>Placebo or no treatment (usual care)</li> <li>Other active comparator listed above, including combinations</li> </ul>
Outcome	Primary outcomes: All-cause mortality at ≥12 months; cardiac mortality at ≥12 months; health-related quality of life at 6 months and ≥12 months; onset of symptoms or progression in NYHA class at ≥12 months; and need for heart valve intervention (surgical or transcatheter) at ≥12 months Secondary outcomes: Exercise tolerance and withdrawal from the trial due to adverse events at 6 and 12 months
Study design	Randomised controlled trial
Timeframe	Long-term
Additional information	None

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#### J.1.160 Research recommendation

**J.1.131** What is the clinical and cost-effectiveness of ACE inhibitors, beta-blockers and diuretics for adults with primary severe mitral regurgitation?

8

# J.1.192 Why this is important

- Patients with primary severe mitral regurgitation may develop symptoms of heart failure and
- may be unsuitable for an intervention or waiting for it to be performed. Whilst ACE inhibitors,

- beta-blockers and diuretics have proven benefit in patients with heart failure due to reduced
   left ventricular systolic function, there is no robust evidence for their effectiveness in patients
- 3 with primary severe mitral regurgitation.

5 Perceived benefit in delaying the need for intervention in patients with primary severe mitral

6 regurgitation has not been demonstrated and there has been some evidence of harm.

### J.1.13 Rationale for research recommendation

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Robust evidence covering this area would allow improved management of those with primary severe mitral regurgitation.
There was very limited evidence on ACE inhibitors, beta- blockers and diuretics for mitral regurgitation. Additional evidence may enable recommendations to be made in the future.
Robust evidence may inform the use of ACE inhibitors, beta-blockers and diuretics in the primary severe mitral regurgitation population as there is currently limited evidence to support this.
None known
There was insufficient evidence to draw conclusions about the relative benefits and harms of ACE-I, beta-blockers and diuretics based on the evidence available. The evidence was based on a small number of studies with a small number of participants. The populations in these studies was younger than that which would be seen on average in the UK, and so may not be representative.
Patients with primary severe mitral regurgitation and heart failure symptoms that are unsuitable for intervention are a subgroup of patients with primary severe mitral regurgitation that needs to be addressed. They also have certain differences compared to the general heart failure population.

# J.1.104 Modified PICO table

11

Population	Inclusion Adults aged 18 years and over with diagnosed primary severe mitral regurgitation (asymptomatic with no indication for intervention or symptomatic and unsuitable for intervention)
Intervention	<ul> <li>Angiotensin-converting enzyme (ACE) inhibitors</li> <li>Beta blockers</li> <li>Diuretics</li> </ul>

	<ul> <li>Any combination of 2 or more of the above</li> </ul>
Comparator	<ul> <li>Placebo or no treatment (usual care)</li> <li>Other active comparator listed above, including combinations</li> </ul>
Outcome	Primary outcomes: All-cause mortality at ≥12 months; cardiac mortality at ≥12 months; health-related quality of life at 6 months and ≥12 months; onset of symptoms or progression in NYHA class at ≥12 months; and need for heart valve intervention (surgical or transcatheter) at ≥12 months Secondary outcomes: Exercise tolerance and withdrawal from the trial due to adverse events at 6 and 12 months
Study design	Randomised controlled trial
Timeframe	Long-term
Additional information	None

# J.2 Heart failure and concomitant heart valve disease

4

#### J.25 Research recommendation

**J.262** What is the clinical and cost effectiveness of beta-blockers for adults ≥75 years with non-rheumatic/calcific mitral stenosis, in both sinus rhythm and atrial fibrillation?

#### J.28 Why this is important

- 9 Calcific mitral stenosis is an increasing in prevalence heart valve disease in the UK with
- ageing of the population. Surgical mitral valve replacement in these patients carries a higher
- 11 risk than other heart valve procedures for the same individual because of technical
- particularities of the procedure and morphologic aspects of the disease. Furthermore, current
- evidence on transcatheter valve implantation in mitral annular calcification is limited and
- 14 suggests that this procedure has only a compassionate role. Consequently, it is important to
- provide pharmacological management of symptoms, classically offered for several decades
- in individuals with rheumatic mitral stenosis. Slowing down the heart rate reduces the
- 17 pressure gradient through the stenotic mitral valve, as demonstrated in classic studies of
- heart catheterisation and known from clinical practice on echocardiography. This was always
- 19 thought to result in an improvement in symptoms and was the base of pharmacological
- 20 management of mitral stenosis. Most patients with mitral stenosis significant enough to
- 21 provoke symptoms are in atrial fibrillations, however occasionally patients may have
- 22 preserved sinus rhythm.

# J.2.4 Rationale for research recommendation

2

Importance to 'patients' or the population	Satisfactory management of symptoms may help delay or avoid need for intervention on the valve.
Relevance to NICE guidance	Only 3 out of 6 of the studies in the evidence review specified whether the participants were in sinus rhythm or atrial fibrillation and none included older adults with calcific heart valve disease. Therefore, to inform future updates of this guidance further research is needed on beta blockers, the key drug intervention in this group, for older adults with non-rheumatic/calcific mitral stenosis, in both sinus rhythm and atrial fibrillation. This is to encourage research to clarify whether this form of pharmacological management is safe and effective in the population most relevant to UK clinical practice and for those in both sinus rhythm and atrial fibrillation as there is currently no randomised evidence to answer these important clinical questions.
Relevance to the NHS	Avoid the cost and the risk related to intervention in these patients, improve their quality of life and reduce their need for hospitalisation.
National priorities	None known
Current evidence base	Only 3 out of 6 of the studies in the evidence review specified whether the participants were in sinus rhythm or atrial fibrillation and none included older adults with calcific heart valve disease. It was also noted that a study showing a benefit included only people in sinus rhythm, while the others did not report the numbers in sinus rhythm or atrial fibrillation. This highlights the need to address whether beta blockers are effective in both sinus rhythm and atrial fibrillation.
Equality considerations	Standardisation of care

3

# J.245 Modified PICO table

Population	Inclusion Adults ≥75 years with non-rheumatic/calcific mitral stenosis, in both sinus rhythm and atrial fibrillation
	Exclusion •
	<ul> <li>People with congenital heart valve disease, except bicuspid aortic valve disease. Patients with contraindication to beta-blockers</li> </ul>

Intervention	Beta blockers
Comparator	<ul> <li>Placebo or no treatment</li> <li>Usual care (e.g. following standard heart failure guidelines)</li> </ul>
Outcome	Primary outcomes  All-cause mortality at 12 months; Cardiac mortality at 12 months; Hospital admission due to heart failure at 12 months; Health-related quality of life at 6 months and 12 months  Secondary outcomes  Exercise tolerance; Need for heart valve intervention (surgical or transcatheter) within 12 months; Withdrawal from the study due to adverse events at 6 months and 12 months
Study design	Randomised controlled trial
Timeframe	Long term
Additional information	None

2

#### J.236 Research recommendation

- **J.247** What is the clinical and cost effectiveness of pharmacological management of heart failure
  - for adults with heart failure and severe aortic stenosis, severe aortic regurgitation or severe
  - 6 mitral regurgitation?

## J.278 Why this is important

- 8 Pharmacological management of heart failure may have particularities in patients with aortic
- 9 stenosis or aortic regurgitation or mitral regurgitation, because of particularities of the valve
- 10 disease impact on haemodynamics. For example, in aortic regurgitation, betablockers are
- thought to have a negative impact if they reduce the heart rate significantly, because slower
- heart rate is associated with longer diastole so longer time for a diastolic phenomenon like
- aortic regurgitation to occur and consequently higher regurgitant volume worsening rather
- than improvement heart failure in these patients. Also for example in aortic stenosis,
- peripheral vasodilation for example with new introduction of an ACE-inhibitor in a patient with
- decompensated aortic stenosis and consequent heart failure may worsen haemodynamics
- and lead to cardiogenic shock. Regarding mitral regurgitation, is it more likely that standard
- heart failure management established for heart failure due to reduced left ventricular systolic
- 19 function has no contraindication, but it not known if it has the same beneficial effect as in
- 20 heart failure due to reduced left ventricular systolic function.

21

### J.229 Rationale for research recommendation

Importance to 'patients' or the population	Establish indication and contraindication for pharmacological management agents in this population.
	F - F

Relevance to NICE guidance	For aortic stenosis evidence was available for the ACE-I versus placebo and ARB versus placebo only but was not considered sufficient to base recommendations on. No relevant randomised controlled trials for aortic or primary mitral regurgitation were identified. This meant that there was only limited evidence to determine whether pharmacological interventions improve outcomes for these people.
Relevance to the NHS	Improve standardisation of care, as current practice varies due to uncertainties and lack of evidence.
National priorities	Not known
Current evidence base	There was insufficient evidence to draw conclusions about the relative benefits and harms of ACE-I or ARB compared with placebo and no other comparisons were available for this stratum. Further evidence is needed in order to consider making strong recommendations. No randomised control trials were identified on pharmacological interventions for people with aortic or mitral regurgitation. As there is a lack of information regarding whether or not any pharmacological interventions improve outcomes evidence is required to be able to make strong recommendations.
Equality considerations	Evidence that would allow NICE to make recommendations that would result in standardisation of care.

# J.2.10 Modified PICO table

3

Population	Inclusion  Adults aged 18 years and over with diagnosed heart failure and severe heart valve disease stratified by type:  • aortic [including bicuspid] stenosis  • aortic [including bicuspid] regurgitation  • Primary mitral regurgitation
	<u>Exclusion</u>
	<ul> <li>Pharmacological management in children (17 years and under)</li> </ul>
	<ul> <li>People with congenital heart valve disease, except bicuspid aortic valve disease.</li> </ul>
	<ul> <li>Patients that can have valve intervention immediately after the diagnosis of their condition rather than be treated medically for a period of time.</li> </ul>

Intervention	<ul> <li>Angiotensin-converting enzyme (ACE) inhibitors</li> <li>Angiotensin-II receptor blockers (ARBs)</li> <li>Beta blockers</li> <li>Digoxin</li> <li>Diuretics</li> <li>Nitrates (including nitroprusside)</li> <li>Any combination of 2 or more of the above</li> </ul>
Comparator	<ul> <li>Placebo or no treatment</li> <li>Usual care (e.g. following standard heart failure guidelines: ACE + beta-blocker + diuretic)</li> <li>Other active comparator listed above, including combinations</li> </ul>
Outcome	Primary outcomes: All-cause mortality at 12 months; Cardiac mortality at 12 months; Hospital admission due to heart failure at 12 months; Health-related quality of life at 6 months and 12 months and Exercise tolerance  Secondary outcomes: Need for heart valve intervention (surgical or transcatheter) within 12 months; and withdrawal from the study due to adverse events at 6 months and 12 months
Study design	Randomised controlled trial
Timeframe	Long term
Additional information	None