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**Is there evidence for statins in the treatment of aortic valve stenosis?**

Akin I *et al*. Statins in aortic stenosis

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**Abstract**

Research revealed that the pathogenesis of aortic stenosis (AS) not merely comprises of a mechanical wear and tear process yet that active biological processes, similar to those of coronary artery disease (CAD) are involved, a promising role for statins in disease-modifying therapy was suggested. However, recently, many prospective studies could not observe decreased progression nor regression of the disease. Here, we review the current knowledge on the pathomechanisms of AS and its similarities and differences with atherosclerosis. Moreover, we discuss whether there is still a place for statins in the treatment of particular AS patient subgroups.

**Key words:** Aortic stenosis; Statins; HMG-CoA enzyme inhibitor; Coronary artery stenosis; Aortic valve surgery

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**Core tip:** Aortic stenosis is a age-dependent and growing disease. As there are several similarities with atherosclerotic disease of other regions there are growing research on underlying pathopyhsiology. The treatment benefit of classic atherosclerosis treatment is evaluated in case of aortic valve stenosis.

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**INTRODUCTION**

Aortic stenosis (AS) is the most common heart valve disease in Western countries. It affects 12.4 % of the population over the age of 75 years[1] with a male predominance of 80 % in symptomatic AS[2]. AS is usually characterized by an asymptomatic latent period of several years or even decades before patients suffer from discomforts. However, once patients with severe AS display symptoms (typically syncope, angina and/or dyspnoea), the prognosis is poor with a 2- and 5-year survival rate of 50 and 20 %, respectively; hence valve replacement surgery is usually mandated[3]. Since the elderly population in Western countries is expected to double by 2050, it is imperative to define early diagnostic and treatment strategies. Even though AS or the fibrous thickening and calcification of the valves is a widespread disease, its underlying molecular mechanisms are still unknown[4]. Traditionally, it was accepted that AS is caused by the progression of calcium deposition continued to aortic valvular leaflets. However, research demonstrating the involvement of active cellular processes over the past 15 years[5]. Genetic polymorphism seems to influence the degree of aortic valve sclerosis[6], with a focus on chronic inflammatory processes. Moreover, it has become clear that the development of AS shows many similarities with atherosclerosis including infiltration and retention of lipoproteins, lipids, T-lymphocyte and other inflammatory cells, as well as osteoblastic activation[7-9]. Accordingly, numerous studies demonstrated a strong associated to coronary artery disease (CAD) and many of its risk factors, including hypercholesterolemia[10,11]. This opened the field to develop effective disease-modifying strategies, which would halt or even regress the disease, thus avoiding the need of surgical or interventional replacement. At first, statins seem to be the most obvious medical treatment choice since its use is well-established for the primary and secondary prevention of CAD[12]. Moreover, statins have been shown to exert pleiotropic effects beyond their cholesterol-lowering effect, such as anti-oxidation, plaques stabilization and reduction of vascular inflammatory processes[13,14]. Numerous retrospective observational studies showed a delay in AS progression when statins were administered concomitantly which encouraged the use of statins for treatment of AS[15-19]. However, 3 recent large-scale prospective randomized clinical trials investigating the effects of intense lipid-lowering therapy with statins showed no effect on neither progression, nor regression of AS. Meanwhile, new insights are emerging, demonstrating distinct differences in (molecular) pathology between CAD and AS. Here, we review what is known today about the pathogenesis of AS and the potential influence of statin therapy in AS patients, with a distinction between degenerative or calcific AS, congenital AS, AS with coronary heart disease as comorbidity, and, treatment before and after aortic valve surgery.

**PATHOMECHANISMS OF DEGENERATIVE OR CALCIFIC AS AND CAD: SIMILARITIES AND DIFFERENCES**

Since it became clear that AS is not merely the result of mechanical stress and ageing, but rather an active biological processes involving inflammatory contributing to the disease; pharmacological treatment possibilities were completely opposed to valve replacement surgery inevitable at the time when AS has reached a severe symptomatic status. What has been revealed of the current knowledge of the pathogenesis of AS so far, revealed many similarities with CAD. Hence, it was obvious to turn to statins for the treatment of AS since they have proven efficacy for the primary and secondary prevention of CAD. Nevertheless, it is shown that, besides lipid-lowering, statins display pleiotropic effects such as anti-inflammatory and antioxidant effects as well as plaque-stabilization and improvement of endothelium dysfunction[20] (Figure 1). The potential use of statins was initially encouraged by positive results from numerous retrospective studies. Yet recently, three prospective randomized studies showed neither regression, nor reduction in progression of AS: SEAS (Simvastatin and Ezetimibe in Aortic Stenosis), SALTIRE (Scottish Aortic Stenosis and Lipid Lowering Trial, Impact on Regression) and ASTRONOMER (Aortic Stenosis Progression Observation: Measuring Effects of Rosuvastatin)[21-23]. All three trials demonstrated that extensive lipid-lowering induced by statins failed to correlate with neither improvement of aortic-jet velocity of the valve, nor with regression of valvular calcification. Such disappointing results were surprising because of the many well recognized similarities between CAD and AS in terms of pathogenesis. Moreover, statin use has become an established principle for CAD. Furthermore, similar risk factors have been identified for both diseases such as age, sex (male gender), dyslipidaemia, hypertension, smoking, diabetes and renal dysfunction[24,25]. Aortic valve stenosis can be characterized an ‘early lesion’, which are similar to those of atherosclerotic plaques in vessels[8] and both CAD and AS are characterized by infiltration and accumulation of (oxidized) low-density lipoproteins (LDL) and T-lymphocytes, thus filling tissue inflammation and the release of pro-inflammatory cytokines such as tumour necrosis factor (TNF)-α and interleukin (IL)-1α which in turn induce cell proliferation[7,9,26]. Calcification and osteogenesis, mediated by inflammation and processes involving metalloproteinases, have been identified as players in the pathogenesis of AS[27-29]. Finally, neo-angiogenesis contributes to AS development[30,31] and is responsible for reduced concentrations of the anti-angiogenesis protein chondromodulin-1 in damaged aortic valve tissue[32]. The resulting cell apoptosis, extracellular matrix formation and consequent thickening and calcification of the cusps, decreases aortic valve mobility and orifice areas, ultimately leading to an increased pressure gradient[33]. Nevertheless, there are some fundamental differences in the underlying molecular mechanisms with an early inflammation affecting fibroblasts in AS as opposed to late onset of inflammation in smooth muscle cells in CAD. Yet, plaque instability is not the leading clinical problem with AS; however clearly the main cause of symptoms with CAD[34]. It is important to note that those retrospective studies on the effect of statins in AS included patients already exposed to statins prescribed for primary or secondary prevention of CAD. Moreover, study patients had only mild to moderate AS as severe AS was excluded. Based on sub-analyses of the 3 RCT’s, several research groups support the lipid hypothesis as a common denominator of both diseases, with similarities between atherosclerosis and AS seen in initial stages of AS. Later on, disease progression is mostly determined by plasticity and structure of the leaflets, and by mechanical stress[35,36]. Yet, results of a recent meta-analysis challenged suggested role of statins to prevent the onset of AS in non-symptomatic patients[37]. They pooled high risk patients without known AS from 3 large scale RCT’s who evaluated the effect of high (80 mg) and normal (10 mg) daily doses of atorvastatin and found no significant differences between placebo, low or high atorvastatin with regards to the onset of AS. Furthermore, across the board any subtle correlation between hyperlipidaemia and AS progression[15,17] turned out to be inconsistent[18,19].

**Effect of Statins on AS in patients with CAD**

As much as 40 % of patients undergoing aortic valve replacement surgery suffers from atherosclerosis as well. Moreover, AS progresses faster in older patients with CAD[2]. There is evidence of beneficial effects of statins in presence of CAD[38]. Dyslipidaemia is an independent predictor of AS progression and adequate lipid-lowering by statin use in patients with CAD has beneficial effects on valve integrity[39]. Yet, a quantitative relationship between lipid-lowering and changes in AS progression has not been found. It is worth mentioning that most studies investigated patients with intermediate or advanced AS which appears difficult to modify by drugs; however, a potential effect in early AS could still be debated[40]. Another point to address is patient selection bias, especially in the SEAS trial that excluded patients with diabetes or CAD. Even though no correlation between statin use and beneficial effects on AS have been shown, it is highly likely that patients with AS and relevant comorbidities such as atherosclerosis, which is a common clinical scenario, may be positively affected[41,42]. Indeed, statins would mitigate AS progression only when hyperlipidaemia is present[43] but more profound research is needed. Current guidelines suggest offering patients with comorbidities such as CAD, diabetes or hyperlipidaemia statins. Conversely, untreated metabolic syndrome in patients with AS is related to faster stenosis progression and worse prognosis[44]. These observations were more pronounced in younger patients (< 57 years of age). Interestingly, however, statins in younger patients with metabolic syndrome and AS turned out to be disadvantageous with AS progression and lower insulin resistance supporting the notion that lipophilic statins may actually induce diabetes type II[45,46]; this effect is even more pronounced when patients suffer from metabolic syndrome as well[47]. This effect was not seen in older patients, suggesting that the pathophysiological mechanisms of AS progression may be related to the patient’s age[44].

**CONGENITAL AORTIC STENOSIS**

Congenital defects resulting in AS include partially fused leaflets, thickened leaflets, narrowing of the supravalvular or subvalvular area, and, most frequently, bicuspidity. A biscupid aortic valve is the result of the fusion of two of the three leaflets during the developmental phase and is with an incidence of almost 2% the most common congenital cardiac malformation[48]. The abnormal structure of the bicuspid aortic valve results in excess stress onto leaflets, resulting in valvular thickening, calcification, and increased rigidity and restricted aortic orifice. For that reason, AS is the most frequent complication of bicuspid aortic valve. While randomized prospective trials mainly included older AS patients, the PROCAS (Progression of Stenosis in Adult Patients with Congenital Aortic Stenosis) trial was designed to evaluate the effect of statins on the progression of stenosis in young asymptomatic adults (aged between 18 and 45) with congenital AS. Not surprising, the investigators did not find any benefit from extensive lipid-lowering by 10 mg rosuvastatin, independent of the degree of valve calcification[49]. These findings are in line with those of the bicuspid valve patients subgroup of the ASTRONOMER trial[23].

**STATIN THERAPY AND AORTIC VALVE SURGERY**

Because of their anti-ischemic and anti-inflammatory effects and protection of the endothelium statins have been suggested to reduce atrial fibrillation after aortic valve surgery, yet with conflicting[50,51]. Some beneficial effects of perioperative statin therapy on mortality, stroke, renal insufficiency, length of ICU and hospital stay were found[52,53] however leading to the conclusion that preoperative statin use is justified in case of coronary artery bypass grafting (CABG). Indeed, advantageous effects on the above-mentioned endpoints have been demonstrated in only CABG patients and not in others[54-56]. Conversely, just to add to the confusion statin were also reported to delay restenosis after balloon aortic valvuloplasty or bioprosthetic valve replacement[57-59], but findings were not[60]. With pre- and postoperative statin therapy in patients undergoing aortic valve replacement an increased long-term survival was found with biological valve replacement but not with mitral valve repair or mechanical valve replacement[61]. Finally, a retrospective study of bicuspid aortic valve patients undergoing surgery showed a significant decrease in ascending aortic dilatation when exposed to statins preoperatively[62]. Clearly, more clinical data are required to justify perioperative use of statin therapy.

**FUTURE CONSIDERATIONS**

Future research in this context needs to focus both, elucidating the molecular pathways involved in the pathogenesis of AS and developing potential pharmacological treatment strategies. In that regard, it is of interest to determine both vascular and valvular aspects and when they occur during the evolution of AS. As such, different treatment approaches might apply for distinct stages of the disease. In addition, defining predisposing genetic deviations may help define preventative and curative approaches to slow disease progression. Additionally, in the quest for new pharmacological agents to treat AS, low-cost accurate animal model are missing; only ageing swine can develop AS[34]. At present, there is no evidence that statins halt the progression or induce the regression of AS. The notion that their administration is harmless and devoid any side effects is untrue as we know adverse effects of statins in asymptomatic AS patients without concomitant diseases may in fact induce new risk factors (diabetes mellitus, aortic valve calcification)[44,63]. In conclusion, the general consensus to date is that treatment with statins is not recommended in patients with valvular aortic stenosis and in absence of standard indications to lipid-lowering treatment.

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**Figure 1 Schematic overview of statin effects on aortic valve calcification.**