Dietary patterns, food groups, and incidence of aortic valve stenosis: A prospective cohort study

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**ABSTRACT**

**Background:** The role of diet in the development of aortic valve stenosis (AVS) is unknown. We therefore examined the associations of two dietary patterns, including a modified Dietary Approaches to Stop Hypertension (mDASH) diet and a modified Mediterranean (mMED) diet, and the food items included in these dietary patterns with incidence of aortic valve stenosis (AVS) in a population-based cohort study.

**Methods:** The study cohort comprised 74,401 Swedish adults (54% men) who were free of cardiovascular disease at the time of completion of a baseline questionnaire about habitual diet and other risk factors for chronic diseases. Participants were followed-up through linkage with nationwide registers on hospitalization and causes of death.

**Results:** During 1,132,617 person-years (mean 15.2 years) of follow-up, 1338 incident AVS cases (801 in men and 537 in women) were ascertained. We found no significant associations of the mDASH and mMED dietary patterns or the food groups and beverages included in these diets (i.e., fruit, vegetables, legumes and nuts, whole grains, fish, low-fat dairy foods, full-fat dairy foods, red and processed meat, and sweetened beverages) with risk of AVS. The hazard ratios (95% confidence interval) of AVS per one standard deviation increase in the mDASH and mMED diet scores were respectively 1.02 (0.96–1.07) and 1.00 (0.95–1.06).

**Conclusion:** This study found no evidence that diet plays a role in the development of AVS.

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1. **Introduction**

Aortic valve stenosis (AVS) is a common disease, with a prevalence reaching nearly 10% in populations over age 80 years [1]. The established risk factors for AVS are few, but mounting evidence indicates that cardiometabolic risk factors, such as obesity [2–4], diabetes [2,3,5,6], hypertension [2,3,5,7], and dyslipidemia [5,8–12], increase the risk of AVS. Diet influences cardiometabolic risk factors [13–15] and thus may be implicated in the development of AVS. A healthy diet rich in fruit, vegetables, legumes, low-fat dairy foods, and fish and reduced in red meat, processed meat, and foods rich in added sugars has been shown to be associated with reduced risk of major cardiovascular diseases, including coronary heart disease, heart failure, and stroke [14,16–19]. However, whether a healthy diet influences the risk of AVS is unknown.

The aim of this study was to investigate the associations of two healthy dietary patterns, including the Dietary Approaches to Stop Hypertension diet and the Mediterranean diet, with incidence of AVS in a population-based cohort of Swedish adults. Furthermore, we examined whether individual food groups and beverages included in these two dietary patterns were associated with risk of AVS.

2. **Methods**

2.1. Study population

Data for this study was available from the Swedish Infrastructure for Medical Population-based Life-course Environmental Research (SIMPLER) database. SIMPLER consists of data from the Cohort of Swedish Men, a cohort of 48,850 Swedish men aged 45–79 at enrolment in the late autumn of 1997, and from the Swedish Mammography Cohort, a cohort of 39,227 Swedish women aged 49–83 years in 1997. For this study, we excluded participants with an erroneous or a missing personal identification number (\(n = 540\)); those who died (\(n = 81\)) or had diagnosis of cancer (\(n = 4403\)) or cardiovascular disease (AVS, ischemic heart disease, heart failure, or ischemic stroke; \(n = 7796\)) prior to 1 January 1998; and those with implausibly high or low total energy intake, probably

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reflecting careful completion of the food-frequency questionnaire (n = 856). This left 74,401 individuals (39,926 men and 34,475 women) for the present analysis. The Regional Ethical Review Board at Karolinska Institutet in Stockholm, Sweden, approved the study.

2.2. Assessment of diet and other exposures

At baseline in 1997, participants provided information on educational attainment, habitual diet, alcohol consumption, smoking status and history, weight, height and history of diabetes, hypertension, and hypercholesterolemia. Diet was assessed using a validated food frequency questionnaire [20]. Participants were asked to indicate their average consumption of 94 food items over the past year. For most items, eight predefined frequency categories ranging from never to ≥ 3 times/week were provided. For commonly consumed foods (e.g., milk, bread, and sweetened beverages), participants were asked to indicate their exact consumption per day or per week. We constructed two healthy dietary patterns based on modified versions of the Dietary Approaches to Stop Hypertension (mDASH) and Mediterranean (mMED) diets, as described in detail previously [18,21]. The dietary patterns were slightly modified to better match the Swedish diet. In the present study, the mDASH diet included fruits, vegetables, legumes and nuts, whole grains, and low-fat dairy foods, which were considered healthy foods, as well as red meat and processed meat and sweetened beverages, which were considered less healthy foods. The mMED diet consisted of fruits, vegetables, legumes and nuts, and whole grains. It further included fish, olive oil use and low to moderate alcohol consumption as beneficial components, and full-fat dairy foods and red meat and processed meat as less healthy foods. We grouped participants into sex-specific quintiles according to their consumption of each food group. For the healthy food groups, participants were assigned a score from 1 to 5 from the lowest to the highest quintile. Scores were reversed for the less healthy foods. The scores were summed to create mDASH and mMED diet scores, with higher scores indicating higher adherence to these healthy dietary patterns.

2.3. Ascertainment of AVS cases and follow-up

The personal identification number assigned to each Swedish resident was used for linkage of participants to the Swedish National Patient Register and Swedish Cause of Death Register. Incident cases of AVS were identified using the International Classification of Diseases 10th revision codes I35.0 and I35.2. Participants were followed up from January 1, 1998 until the date of diagnosis of AVS, date of death, or December 31, 2014, whichever was earlier.

2.4. Statistical analysis

Analyses were conducted using Cox proportional hazards regression models, with age as the time scale and stratified by sex. All multivariable models included adjustment for education (less than high school, high school, or university), total energy intake (continuous), alcohol consumption (never drinker, former drinker, or current drinker of < 1, 1–6, 7–14, 14–21, and > 21 drinks/week) (only in analyses of the mDASH diet and food groups since the mMED diet included alcohol intake), and smoking (never, former -20 pack-years, former ≥ 20 pack-years, current -20 pack-years, or current ≥ 20 pack-years). In a second (main) multivariable model, we further adjusted for potential intermediates, including body mass index (weight divided by the square of height; < 22.5, 22.5–24.9, 25.0–29.9, or ≥ 30 kg/m²) and a history of diabetes, hypertension, and hypercholesterolemia. Adjustment for physical activity (exercise and walking/bicycling) did not alter the results. Physical activity was therefore not included in the final multivariable models.

The main results are presented per one standard deviation increment of the mDASH and mMED dietary patterns and food groups. In secondary analyses, we categorized participants into quintiles according to the mDASH and mMED diet scores. In a sensitivity analysis, we excluded participants with a history of diabetes, hypertension, or hypercholesterolemia at baseline who may have changed their dietary behaviors. To correct for multiple testing, we imposed a Bonferroni-corrected significance threshold of 0.005 (i.e., 0.05/11 dietary factors examined). All analyses were performed using SAS, version 9.4 (SAS Institute, Cary, NC).

3. Results

Participants with high adherence to the mDASH and mMED dietary patterns were slightly leaner, were more likely to have a postsecondary

Table 1: Baseline characteristics by quintiles of the mDASH and mMED dietary patterns.

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>mDASH dietary pattern</th>
<th>mMED dietary pattern</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Q1 (n = 16,809)</td>
<td>Q2 (n = 15,029)</td>
</tr>
<tr>
<td></td>
<td>Q1 (n = 15,444)</td>
<td>Q2 (n = 16,167)</td>
</tr>
<tr>
<td>Age, years</td>
<td>60.5 ± 9.6</td>
<td>59.9 ± 9.4</td>
</tr>
<tr>
<td></td>
<td>60.5 ± 9.8</td>
<td>60.5 ± 9.5</td>
</tr>
<tr>
<td>Body mass index, kg/m²</td>
<td>25.3 ± 3.9</td>
<td>25.5 ± 3.7</td>
</tr>
<tr>
<td></td>
<td>25.5 ± 3.9</td>
<td>25.5 ± 3.7</td>
</tr>
<tr>
<td>Postsecondary education, %</td>
<td>12.2</td>
<td>15.1</td>
</tr>
<tr>
<td></td>
<td>11.0</td>
<td>14.6</td>
</tr>
<tr>
<td>Current smoker, %</td>
<td>31.1</td>
<td>26.7</td>
</tr>
<tr>
<td></td>
<td>32.6</td>
<td>27.1</td>
</tr>
<tr>
<td>History of diabetes, %</td>
<td>4.1</td>
<td>4.4</td>
</tr>
<tr>
<td></td>
<td>4.4</td>
<td>4.9</td>
</tr>
<tr>
<td>History of hypertension, %</td>
<td>21.2</td>
<td>21.2</td>
</tr>
<tr>
<td></td>
<td>20.3</td>
<td>21.3</td>
</tr>
<tr>
<td>History of hypercholesterolemia, %</td>
<td>9.4</td>
<td>10.3</td>
</tr>
<tr>
<td></td>
<td>9.1</td>
<td>10.3</td>
</tr>
</tbody>
</table>

- Fruit, servings/day
- Vegetables, servings/day
- Legumes and nuts, servings/day
- Whole grains, servings/day
- Fish, servings/day
- Low-fat dairy, servings/day
- Full-fat dairy, servings/day
- Red and processed meat, servings/day
- Sweetened beverages, servings/day
- Alcohol intake, drinks/wk
- Olive oil use, %
- Macronutrients

a Age-standardized to the age distribution of the study population at baseline. Values are means ± standard deviation if not otherwise indicated.

b Component of the mDASH diet.
cc Component of the mMED diet.
d Among current drinkers.

Adjusted for total energy intake.

We refer you to the full text for a detailed interpretation of the results. Please cite this article as: S.C. Larsson, A. Wolk and M. Bäck, Dietary patterns, food groups, and incidence of aortic valve stenosis: A prospective cohort study, International Journal of Cardiology, https://doi.org/10.1016/j.ijcard.2018.11.007.
education and a history of diabetes and hypercholesterolemia, and were
less likely to be current smoker compared with those with low adher-
ence to these dietary patterns (Table 1). Participants with high adher-
ence to the mDASH dietary pattern were somewhat less likely to have
a history of hypertension, whereas those with high adherence to the
mMED dietary pattern were more likely to have a history of hyperten-
sion compared with low adherence to each of these dietary patterns
(Table 1). High adherence to the two healthy dietary patterns was asso-
ciated with lower total fat intake and higher carbohydrate and protein
intake (Table 1).

During 1,132,617 person-years (mean 15.2 years) of follow-up, 1,338
incident AVS cases (801 in men and 537 in women) were ascertained.
The mDASH and mMED dietary patterns were not significantly asso-
ciated with incidence of AVS (Fig. 1 and Table 2). The hazard ratios (95% confi-
dence interval) per one standard deviation increment of the mDASH and
mMED dietary patterns were respectively 1.02 (0.96–1.07) and 0.94 (0.87–1.06)
(Fig. 1). In a sensitivity analysis excluding participants with a
history of diabetes, hypertension, or hypercholesterolemia at baseline,
the corresponding hazard ratios (95% confidence interval) were 0.96
(0.89–1.03) for the mDASH diet and 0.94 (0.87–1.02) for the mMED diet.

### Table 2

<table>
<thead>
<tr>
<th>Dietary pattern</th>
<th>No. of cases</th>
<th>Person-years</th>
<th>Age- and sex-adjusted HR (95% CI)</th>
<th>Multivariable HR (95% CI)a</th>
<th>Multivariable HR (95% CI)b</th>
</tr>
</thead>
<tbody>
<tr>
<td>mDASH</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Q1</td>
<td>289</td>
<td>250,805</td>
<td>1.00 (reference)</td>
<td>1.00 (reference)</td>
<td>1.00 (reference)</td>
</tr>
<tr>
<td>Q2</td>
<td>293</td>
<td>227,612</td>
<td>1.10 (0.93–1.30)</td>
<td>1.13 (0.96–1.33)</td>
<td>1.13 (0.96–1.31)</td>
</tr>
<tr>
<td>Q3</td>
<td>217</td>
<td>195,446</td>
<td>0.95 (0.79–1.13)</td>
<td>0.99 (0.83–1.19)</td>
<td>0.98 (0.82–1.17)</td>
</tr>
<tr>
<td>Q4</td>
<td>308</td>
<td>255,898</td>
<td>1.03 (0.87–1.20)</td>
<td>1.11 (0.94–1.31)</td>
<td>1.10 (0.93–1.30)</td>
</tr>
<tr>
<td>Q5</td>
<td>231</td>
<td>202,856</td>
<td>0.96 (0.81–1.14)</td>
<td>1.08 (0.90–1.29)</td>
<td>1.06 (0.89–1.27)</td>
</tr>
<tr>
<td>mMED</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Q1</td>
<td>293</td>
<td>228,948</td>
<td>1.00 (reference)</td>
<td>1.00 (reference)</td>
<td>1.00 (reference)</td>
</tr>
<tr>
<td>Q2</td>
<td>292</td>
<td>250,845</td>
<td>0.90 (0.77–1.06)</td>
<td>0.93 (0.79–1.09)</td>
<td>0.92 (0.78–1.08)</td>
</tr>
<tr>
<td>Q3</td>
<td>253</td>
<td>221,230</td>
<td>0.92 (0.78–1.09)</td>
<td>0.97 (0.82–1.15)</td>
<td>0.96 (0.81–1.14)</td>
</tr>
<tr>
<td>Q4</td>
<td>260</td>
<td>217,810</td>
<td>0.96 (0.81–1.14)</td>
<td>1.03 (0.87–1.23)</td>
<td>1.02 (0.86–1.21)</td>
</tr>
<tr>
<td>Q5</td>
<td>240</td>
<td>213,784</td>
<td>0.92 (0.77–1.09)</td>
<td>1.02 (0.86–1.22)</td>
<td>1.00 (0.84–1.20)</td>
</tr>
</tbody>
</table>

HR, hazard ratio; CI, confidence interval; mDASH, modified Dietary Approaches to Stop Hypertension; mMED, modified Mediterranean; Q, quintile.

* Hazard ratios were derived from Cox proportional hazards regression models with age as the time scale, stratified by sex, and adjusted for education (less than high school, high school or university), total energy intake (continuous), alcohol consumption (never drinker, former drinker, or current drinker of <1, 1–6, 7–14, 14–21, and ≥21 drinks/week), total energy intake (continuous), body mass index (BMI) (among the mDASH and mMED diet patterns), smoking (never, former, or current smoker), and history of diabetes, hypertension, or hypercholesterolemia.

b Further adjusted for body mass index (BMI) (≥22.5, 22.5–24.9, 25.0–29.9, or ≥30 kg/m²) and history of diabetes, hypertension, or hypercholesterolemia.

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Among the food groups included in the mDASH and mMED dietary patterns, sweetened beverage consumption was weakly positively associated with AVS risk (P = 0.01) (Fig. 1) but the P value did not achieve the Bonferroni-corrected statistical significance threshold (P < 0.005) imposed to adjust for multiple testing. Furthermore, in a post-hoc analysis in which participants were grouped into quintiles of sweetened beverage consumption, no significant association was observed in any quintile (hazard ratio for the highest versus lowest quintile, 1.13; 95% confidence interval 0.95–1.33). The other food groups (Fig. 1) and olive oil use (hazard ratio 0.97; 95% confidence interval 0.84–1.12) were not associated with AVS risk.

4. Discussion

Findings from this large cohort of Swedish adults provide no evidence that diet plays a role in the development of AVS. Specifically, we observed no associations of overall healthy dietary patterns or major foods groups and sweetened beverages with risk of AVS. In previous studies based on data from this cohort, we have observed inverse associations of the mDASH and mMED dietary patterns with risk of myocardial infarction, heart failure, and stroke [18,19] as well as with all-cause mortality [22,23]. Furthermore, we have observed that fruit, vegetables, low-fat dairy foods, and fish consumption is inversely associated with risk of myocardial infarction, heart failure, stroke, and all-cause mortality whereas high intakes of red meat, processed meat and sweetened beverages are associated with increased risk of these diseases and mortality [24–34]. Components in these foods could possibly influence the risk of AVS by decreasing oxidation and atherogenicity of low-density lipoprotein cholesterol (antioxidant nutrients and phytochemicals in fruit, vegetables, whole grains, legumes, nuts, and olive oil) [35–37]; by reducing blood pressure (antioxidants and minerals in fruit, vegetables, nuts, olive oil, and low-fat dairy foods) [13,38,39] or inflammation (long-chain omega-3 polyunsaturated fatty acids in fish) [40]; by increasing blood pressure (sodium in for example processed meat) [41]; or by inducing postprandial hyperglycemia, hyperinsulinemia, and obesity (sucrose in sugar-sweetened beverages) [14]. Experimental studies in mice have shown that high-fat diets induce aortic valve disease [42,43]. The mDASH and mMED diets were inversely associated with dietary fat intake in this study but none of the dietary patterns or any of the examined food groups were associated with incidence of AVS. In previous studies assessing the risk factors for AVS in this study population we observed no association with nutrient consumption specifically [44], but found an inverse association with light alcohol consumption [45]. To the best of our knowledge, no other epimiologic study has assessed the association of any dietary pattern or specific foods or food groups with risk of AVS.

Important strengths of this study include the large number of incident AVS cases and the objective ascertainment of cases through linkage to nationwide population-based registers. A limitation is the use of a self-administered food-frequency questionnaire to assess dietary intake. In addition, only a baseline measurement of diet was used and participants may have changed their diet during follow-up. Thus, some degree of measurement error was inevitable. Nevertheless, the validity of the dietary questionnaire is reasonably good [20] and we have previously observed associations between diet and many other chronic diseases and mortality in this study population (e.g., [18,19,21–34]). This suggests that this dietary questionnaire can detect associations between diet and disease in this study population and that lack of association of the healthy dietary patterns and major foods groups with AVS likely represent a true lack of association. However, we cannot rule out that we may have overlooked a weak association. Another limitation is the observational design. We therefore cannot entirely rule out the possibility that the lack of association between diet and AVS is due to residual confounding. Another potential shortcoming is that despite high validity of the diagnoses in the Swedish National Patient Register [46], AVS incidence may be slightly underestimated. The generalizability of our results to individuals not seeking specialist care is thus unknown.

In conclusion, this study found no evidence that diet plays a role in the development of AVS. Since this is, to the best of our knowledge, the first epidemiologic study evaluating the association between diet and AVS risk, further large cohort studies on dietary patterns, foods, and specific dietary factors (e.g., nutrients and other bioactive dietary compounds) in relation to incidence of AVS may be warranted.

Funding

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Conflict of interest

The authors have no conflicts of interest to declare.

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