Uveitis as a Risk Factor for Developing Acute Myocardial Infarction in Ankylosing Spondylitis: A National Population-Based Longitudinal Cohort Study

Yi-Fen Lai1, Ting-Yi Lin1, Wu-Chien Chien2,3,4, Chien-An Sun5,6, Chi-Hsiang Chung2,3,7, Yi-Hao Chen1, Jiann-Torng Chen1 and Ching-Long Chen1*

1 Department of Ophthalmology, Tri-Service General Hospital, National Defense Medical Center, Taipei, Taiwan, 2 Department of Medical Research, Tri-Service General Hospital, National Defense Medical Center, Taipei, Taiwan, 3 School of Public Health, National Defense Medical Center, Taipei, Taiwan, 4 Graduate of Life Sciences, National Defense Medical Center, Taipei, Taiwan, 5 Department of Public Health, College of Medicine, Fu-Jen Catholic University, New Taipei City, Taiwan, 6 Big Data Research Center, College of Medicine, Fu-Jen Catholic University, New Taipei City, Taiwan, 7 Taiwanese Injury Prevention and Safety Promotion Association, Taipei, Taiwan

Background: Ankylosing spondylitis (AS) is a chronic inflammatory disease. Excess cardiovascular risks were well recognized in patients with AS and were attributed to prolonged systemic inflammation. Uveitis is one of the most common extra-articular symptoms of AS and is also considered an indicator of systemic inflammation. This study aimed to investigate whether uveitis was a risk factor for developing acute myocardial infarction (AMI) in patients with AS using the National Health Insurance Research Database (NHIRD).

Methods: Data were collected from the NHIRD over a fifteen-year period. Variables were analyzed using the Pearson chi-square test and Fisher’s exact test. Risk factors for the occurrence of AMI were examined by calculating hazard ratio. Kaplan-Meier analysis was performed to compare the cumulative incidence of AMI in the uveitis and non-uveitis cohorts.

Results: A total of 5905 patients with AS were enrolled, including 1181 patients with uveitis (20%) and 4724 patients without uveitis (80%). The Kaplan–Meier method with the log-rank test showed that the uveitis group had a significantly higher cumulative hazard for patients with AMI than the non-uveitis group (p < 0.001). The adjusted hazard ratio (aHR) of AMI was higher in the uveitis group than in the non-uveitis group (aHR = 1.653, p < 0.001). Stratified analysis revealed that patients with uveitis had an increased risk of developing AMI regardless of their sex (male/female aHR = 1.688/1.608, p < 0.001). Patients with uveitis in all age groups were independently associated with an increased risk of developing AMI compared to those without uveitis (20–39 years/40–59 years/≥ 60 years, aHR = 1.550, 1.579, 3.240, p < 0.001). Patients with uveitis had a higher probability
INTRODUCTION

Ankylosing spondylitis (AS) is a chronic inflammatory disease that predominantly affects the spine, with a peak onset between the ages of 20 years and 40 years (1). More than 85% of these patients have a specific human leukocyte antigen, known as the HLA-B27 antigen (2). Patients may also experience extra-articular manifestations, such as uveitis, psoriasis, and inflammatory bowel disease. Vascular involvement manifests as aortitis, perioartitis, and aortic valve regurgitation (3). Coronary artery involvement is extremely rare in AS, which is serious and life-threatening when acute myocardial infarction (AMI) occurs (4). As for many systemic inflammatory diseases, AS with excess cardiovascular mortality and morbidity has been well recognized in previous studies (5, 6). The underlying mechanism of AMI may be attributed to the systemic inflammation associated with AS. Prolonged inflammation-induced vascular endothelial injury may result in atherosclerosis and thrombosis, leading to AMI (7).

Uveitis is one of the most common extra-articular symptoms of AS. The prevalence of uveitis in AS is approximately 20–30% (8). Currently, uveitis is believed to be not only an intraocular inflammation but also an indicator of systemic inflammation. Previous studies have demonstrated that the levels of pro-inflammatory cytokines increase in tears, aqueous humor, and serum in active disease (9–11). Uveitis might reflect the severity of systemic inflammation in AS, since inflammation may cause more cytokine cascades toward coronary vascular endothelial injuries, uveitis might be a risk factor for the development of AMI in AS. However, no study has proved this hypothesis.

As AMI is associated with life-threatening consequences (especially in young adults), it is important to know how AS and AMI are associated, including the risk factors of AMI in AS. However, most studies on this subject are case reports or case series, and the lack of a large sample size to investigate the association. Therefore, we used the National Health Insurance Research Database (NHIRD), which contains a comprehensive longitudinal medical record from 23 million people in Taiwan. This approach resulted in a large number and varying features of patients with AS compared to previous studies. This study aimed to investigate the factors that determine the incidence of AMI in AS. In addition, the study elucidated whether uveitis could be a risk factor for the development of AMI in AS. Our ultimate goal is to raise the awareness of physicians about AMI in AS patients with uveitis and provide information to all physicians for multidisciplinary care.

MATERIAL AND METHODS

Research Database

Beginning in 1995, the National Health Insurance (NHI) program was launched in Taiwan. More than 23 million Taiwanese and non-Taiwanese residents (approximately 99.9% of the total population) were included. The National Health Insurance Research Database (NHIRD) contains comprehensive medical records, including outpatients, inpatients, diagnoses, prescriptions, and procedures, which are registered by the International Classification of Diseases, Ninth Revision, Clinical Modification (ICD-9-CM) codes. The ICD-9-CM codes used in this study are listed in Table S1. These data eliminate personal identification and are available in an electronic format for clinicians to conduct statistical analysis.

Study Participants

We conducted a retrospective cohort study between January 2000 and December 2015. All included patients had at least one inpatient claim or outpatient visits more than three times with a diagnosis of AS according to the ICD-9-CM code (n = 21,846, ICD-9-CM code 720.0). Patients younger than 20 years old, with unknown sex, or without tracking were excluded. Patients diagnosed with AS and AMI before the index date were also excluded to ensure that patients had a new onset of AS and new onset of AMI after AS diagnosis. The index date was defined as the date of the first AS diagnosis. The study population was divided into the uveitis group (n = 1181, study cohort) and non-uveitic group (n = 18815). The group of non-uveitic patients matched four-fold with the uveitic group for sex, age, comorbidities, and index date as the comparison cohort (n = 4724). The uveitis group was divided into the anterior uveitis and posterior segment involvement groups. The populations in both groups did not overlap with each other. The uveitic subgroups and comorbidities extracted by ICD-9-CM codes are listed in Table S1. We replaced the Charlson comorbidity index (CCI) with the CCI_R (CCI excluding AMI, diabetes mellitus [DM], hyperlipidemia, hypertension [HTN], cerebrovascular accident [CVA], congestive heart failure [CHF], chronic obstructive pulmonary disease [COPD], asthma, coronary artery disease [CAD], cardiomegaly...
and metabolic syndrome (MetS)) because the removed diseases were also variables in this study. Patients were followed until AMI occurred (n = 138 in the uveitic group; n = 415 in the non-uveitic group) or until the end of the study period. Figure 1 shows a flow chart of the patient selection process.

**Ethical Consideration**

The study protocol was reviewed and approved by the institutional review board of the Tri-Service General Hospital (TSGH IRB No. B-110-35). Because the identities of all patients in the NHIRD were encrypted before data were released, the requirement for signed informed consent of the included patients was waived.

**Statistical Analysis**

The Pearson chi-square test and Fisher’s exact test were used to evaluate differences in categorical variables such as sex, age group, and comorbidities. The threshold for statistical significance was set at p < 0.05. A Cox proportional hazard model was used to estimate the hazard ratio (HR) for the occurrence of AMI based on each variable in the univariate and multivariate analyses. Considering competing risk, Cox regression with Fine and Gray’s competing risk model was also employed for analysis. Kaplan–Meier analysis was performed to estimate the cumulative incidence of AMI in these two cohorts. All statistical analyses were performed using SPSS software (Version 22.0; SPSS Inc., Chicago, IL, USA).

**RESULTS**

Table 1 shows the demographic and clinical characteristics of the enrolled patients with AS at baseline. A total of 5905 patients

![Flowchart of patient selection from the National Health Insurance Research Database in Taiwan.](image-url)
were enrolled, including 1181 (20%) patients with uveitis (uveitis group) and 4724 (80%) patients without uveitis (non-uveitis group). The non-uveitis group was matched four-fold with the uveitis group with age, sex, comorbidities, and index date. The mean age of the total AS patient was 37.72 ± 18.90 years, with the majority of patients (59.10%) younger than 40 years of age. Men accounted for 55.12% of the study population. For comparison of comorbidities at baseline, the prevalence of diabetes and hypertension was 25.15% and 20.51%, respectively. Those with chronic obstructive pulmonary disease and asthma accounted for 16.27% and 19.14% of the total AS cohort, respectively. Fewer patients (< 7%) had hyperlipidemia, cerebrovascular accident, congestive heart failure, coronary artery disease, cardiomegaly, and metabolic syndrome. At the end of follow-up, the percentages of AMI in the uveitis and non-uveitis groups were 11.69% and 8.78%, respectively (p < 0.001). All-cause mortality, which stands for death caused by any event, was 8.55% and 7.79% in the uveitis and non-uveitis groups, respectively (p = 0.332). The mortality caused by AMI in the uveitis group and non-uveitis group was 0.93% and 0.83%, respectively (p=0.723). The mean follow-up time is 9.91 ± 8.56 (range 0.01–15.99) years for all patients. The average time of AMI occurrence after patients included in the study was 2.90 ± 3.17 (range 0.03–15.82) years in the uveitis group and 3.64 ± 4.20 years (range 0.03–15.88) years in the non-uveitis group (p<0.001).

The cumulative risk of developing AMI in patients with AS was calculated using the Kaplan–Meier method (Figure 2). The results showed that the uveitis group had a significantly higher hazard than the non-uveitis group (log-rank test, p < 0.001).

### TABLE 1 | Patients’ clinical characteristics at baseline.

| Uveitis | Total | Without | P |
|---------|-------|---------|---|
| Variables | n | % | n | % | n | % |
| **Total** | 5,905 | 1,181 | 4,724 | 0.999 |
| Gender | | | | 0.999 |
| Male | 3,255 | 55.12 | 2,604 | 55.12 |
| Female | 2,650 | 44.88 | 2,120 | 44.88 |
| Age (years) (mean ± SD) | 37.72 ± 18.90 | 37.61 ± 18.79 | 37.75 ± 18.97 | 0.868 |
| Age group (yrs) | | | | 0.999 |
| 20-39 | 3,490 | 59.10 | 698 | 59.10 |
| 40-59 | 1,655 | 28.03 | 331 | 28.03 |
| ≥60 | 760 | 12.87 | 152 | 12.87 |
| DM | | | | 0.925 |
| Without | 4,420 | 74.85 | 883 | 74.77 |
| With | 1,485 | 25.15 | 298 | 25.23 |
| Hyperlipidemia | | | | 0.810 |
| Without | 5,540 | 93.82 | 1,109 | 93.90 |
| With | 365 | 6.18 | 72 | 6.10 |
| HTN | | | | 0.913 |
| Without | 4,694 | 79.49 | 940 | 79.59 |
| With | 1,211 | 20.51 | 241 | 20.41 |
| CVA | | | | 0.825 |
| Without | 5,568 | 94.29 | 1,112 | 94.16 |
| With | 337 | 5.71 | 69 | 5.84 |
| CHF | | | | 0.763 |
| Without | 5,730 | 97.04 | 1,145 | 96.95 |
| With | 175 | 2.96 | 36 | 3.05 |
| COPD | | | | 0.881 |
| Without | 4,944 | 83.73 | 987 | 83.57 |
| With | 961 | 16.27 | 194 | 16.43 |
| Asthma | | | | 0.835 |
| Without | 4,775 | 80.86 | 952 | 80.61 |
| With | 1,130 | 19.14 | 229 | 19.39 |
| CAD | | | | 0.962 |
| Without | 5,545 | 93.90 | 1,108 | 93.82 |
| With | 360 | 6.10 | 73 | 6.18 |
| Cardiomegaly | | | | 0.898 |
| Without | 5,845 | 98.98 | 1,170 | 99.07 |
| With | 60 | 1.02 | 11 | 0.93 |
| MetS | | | | 0.880 |
| Without | 5,711 | 96.71 | 1,141 | 96.61 |
| With | 194 | 3.29 | 40 | 3.39 |
| CCI_R | | | | 0.162 |

DM, diabetes mellitus; HTN, hypertension; CVA, cerebrovascular accident; CHF, congestive heart failure; COPD, chronic obstructive pulmonary disease; CAD, coronary artery disease; MetS, metabolic syndrome; CCI_R, Charlson comorbidity index excluding acute myocardial infarction, diabetes mellitus, hyperlipidemia, hypertension, cerebrovascular accident, congestive heart failure, chronic obstructive pulmonary disease, asthma, coronary artery disease, cardiomegaly and metabolic syndrome.
without Fine and Gray’s competing risk model are shown in Table 2. The adjusted HR (aHR) with competing risk analysis for AMI was 1.653 times greater in the uveitis group than in the non-uveitis group (95% CI 1.480–1.918). Although men were more likely to develop AMI, the result was not statistically significant after adjusting for confounders (aHR 1.179, p = 0.238). AS patients aged over 40 years showed a higher risk of developing AMI (aHR 1.11 in the 40–59 age group, p = 0.036; aHR 1.718 in the ≥ 60 age group, all p < 0.001). The hazard was significantly higher with age. Adjusted HRs showed that AS patients with DM, hyperlipidemia, HTN, CVA, CHF, COPD, asthma, CAD, and metabolic syndrome had a significantly higher risk of developing AMI, except for AS patients with cardiomegaly.

In the stratified analysis comparing patients with and without uveitis (Table 3), the overall incidence of AMI was 1306.54 per 100,000 person-years in the uveitis group and 885.53 per 100,000 person-years in the non-uveitis group. Patients with uveitis have an increased risk of developing AMI regardless of their sex. The aHR was 1.688 in men and 1.608 in women (p < 0.001). Patients with uveitis in all age groups were independently associated with an increased risk of developing AMI than patients without uveitis, with an aHR of 1.550 in the 20–39 age group, 1.579 in the 40–59 age group, and 3.240 in the ≥ 60 age group (all p < 0.001). The aHR increased with age. For comorbidities, the uveitis group without DM, hyperlipidemia, HTN, CVA, CHF, COPD, asthma, CAD, cardiomegaly, and metabolic syndrome had an increased risk of developing AMI with aHRs of 1.550, 1.581, 1.439, 1.603, 1.659, 1.627, 1.597, 1.577, 1.650, and 1.615, respectively (all p < 0.001). The aHR was higher in the uveitis group with comorbidities than in the uveitis group without comorbidities. The uveitis group with DM, hyperlipidemia, HTN, CVA, CHF, COPD, asthma, CAD, and metabolic syndrome had increased aHR of developing AMI with aHRs of 3.287, 3.102, 4.166, 3.296, 3.261, 3.358, 3.286, 5.187, and 1.932, respectively (all p < 0.001).

The uveitis group (n = 1181) was further divided into the anterior uveitis group (36.6%, n = 432) and posterior segment involvement group (63.4%, n = 749) compared to the non-uveitis group (n = 4724) (Table 4). The overall incidence of AMI was 885.53 per 100,000 person-years in the non-uveitis group, 1,211.31 per 100,000 person-years in the anterior uveitis group, and 1,371.76 person-years in the posterior segment involvement group. The aHR was 1.624 in the anterior uveitis group and 1.673 in the posterior segment involvement group. Uveitis patients with posterior segment involvement had a slightly increased aHR for developing AMI than patients with only anterior uveitis.

**DISCUSSION**

In this population-based cohort study, we enrolled 5905 patients with newly diagnosed AS. Of these patients, 1181 had uveitis and 4724 did not have uveitis. In the mean follow-up of 9.91 years, we found that patients with uveitis had a significantly higher risk (aHR = 1.653) for developing AMI than patients without uveitis. Kaplan–Meier analysis also revealed that the cumulative risk of developing AMI was significantly increased in the uveitis group. In the stratified analysis, the incidence of AMI in the uveitis group was 1306.54 per 100,000 person-years and 1.47-fold higher than that in the non-uveitis group. Uveitis was associated with an increased risk of developing AMI in both sexes, all age groups, patients with and without comorbid DM, hyperlipidemia, HTN, CVA, CHF, COPD, asthma, CAD, and metabolic syndrome. Uveitis patients with comorbidities had a higher adjusted hazard of developing AMI compared to uveitis patients without comorbidities.

The prevalence of AS has geographical differences. AS is more common in Europe and Asia than in Latin America (12).
Previous studies have shown that the prevalence of AS per 10000 is highest in Turkey (49.0), lower in Taiwan (33.7), India (7.0), and lowest in Mexico (2.6) (13–16). The male-to-female ratio in AS also varies across countries, ranging between 1.2 and 7.0 (13, 17); however, these results consistently showed a male predominance. In the present study, the male-to-female ratio was 1.23, which is similar to the results previous study in Asia (18). Previous studies have shown that AS is usually diagnosed in the second or fourth decade of life (19, 20). In our study, more than half of the participants (59.10%) was consistently between 20–40 years of age. In our report, 5.9% of patients with AS had uveitis (1181/19996 = 5.90%). The prevalence of uveitis in AS varies in different studies, ranging from 0% to 30% (8, 21–23) and was even lower (0%) in a Philippines hospital-based study (24).

For comorbidities, the percentages of hypertension and chronic obstructive pulmonary disease were similar to those reported in previous studies (25, 26). The percentage of patients with diabetes and asthma was slightly higher than those reported in previous studies. The percentage of diabetes in a study carried out in the US was reported to be 17.3% in whites and 27.2% in African Americans (27). Asthma patients accounted for 7.3% of a German cohort (28).

Our study population comprised AS patients with uveitis and AS patients without uveitis matched by AS patients with uveitis.  

| Variables | No competing risk in the model | Competing risk in the model |
|-----------|-------------------------------|-----------------------------|
|           | Crude HR 95% CI P Adjusted HR 95% CI P | Crude HR 95% CI P Adjusted HR 95% CI P |
| Uveitis   | Without: Reference 1.821 1.527 2.027 <0.001* | Without: Reference 1.910 1.635 2.067 <0.001* |
|           | With: 1.169 0.790 1.565 0.302 1.144 0.766 1.528 0.309 | With: 1.209 0.856 1.722 0.193 1.179 0.816 1.707 0.238 |
| Gender    | Male: Reference 1.106 1.043 1.308 0.028* | Male: Reference 1.120 1.056 1.400 0.014* |
|           | Female: Reference 1.161 1.205 1.764 <0.001* | Female: Reference 1.737 1.223 1.885 <0.001* |
| Age (yrs) | 20-39: Reference 1.074 1.003 1.266 0.047* | 20-39: Reference 1.120 1.056 1.400 0.014* |
|           | 40-59: Reference 1.511 1.144 1.736 <0.001* | 40-59: Reference 1.718 1.220 1.867 <0.001* |
|           | 60+: Reference 2.947 2.023 4.037 <0.001* | 60+: Reference 3.097 2.168 5.276 <0.001* |
| DM        | Without: Reference 2.586 1.508 4.203 <0.001* | DM: Reference 2.954 1.917 4.991 <0.001* |
|           | With: 2.714 1.560 4.312 <0.001* | With: 2.948 1.817 4.906 <0.001* |
| Hyperlipidemia | Without: Reference 3.063 2.020 5.088 <0.001* | Hyperlipidemia: Reference 3.434 2.492 5.785 <0.001* |
|           | With: 2.157 1.244 2.652 <0.001* | With: 2.284 1.407 2.828 <0.001* |
| HTN       | Without: Reference 1.448 1.039 2.377 0.038* | HTN: Reference 1.551 1.102 2.596 <0.001* |
|           | With: 1.919 1.041 2.606 0.009* | With: 1.940 1.120 2.729 <0.001* |
| CVA       | Without: Reference 2.329 1.470 2.856 <0.001* | CVA: Reference 2.445 1.595 2.985 <0.001* |
|           | With: 2.349 1.532 2.948 <0.001* | With: 2.350 1.535 2.960 <0.001* |
| CHF       | Without: Reference 1.488 1.289 2.704 <0.001* | CHF: Reference 4.334 2.399 7.263 <0.001* |
|           | With: 4.188 2.289 7.094 <0.001* | With: 4.172 2.224 7.033 <0.001* |
| COPD      | Without: Reference 1.000 - - | COPD: Reference 0.000 - - |
|           | With: 2.190 1.294 4.112 <0.001* | With: 2.190 1.294 4.112 <0.001* |
| Asthma    | Without: Reference 0.000 - - | Asthma: Reference 0.000 - - |
|           | With: 1.210 1.144 1.226 <0.001* | With: 1.210 1.144 1.226 <0.001* |
| CAD       | Without: Reference 0.000 - - | CAD: Reference 0.000 - - |
|           | With: 0.000 - - | With: 0.000 - - |
| Cardiomegaly | Without: Reference 0.000 - - | Cardiomegaly: Reference 0.000 - - |
|           | With: 0.000 - - | With: 0.000 - - |
| MetS      | Without: Reference 0.000 - - | MetS: Reference 0.000 - - |
|           | With: 0.000 - - | With: 0.000 - - |

*Indicates statistical significance, p < 0.05. HR, hazard ratio; CI, confidence interval; Adjusted HR, adjusted variables are listed; DM, diabetes mellitus; HTN, hypertension; CVA, cerebrovascular accident; CHF, congestive heart failure; COPD, chronic obstructive pulmonary disease; CAD, coronary artery disease; MetS, metabolic syndrome; Competing risk, All-cause mortality; CCI_R, Charlson comorbidity index excluding acute myocardial infarction, diabetes mellitus, hyperlipidemia, hypertension, cerebrovascular accident, congestive heart failure, chronic obstructive pulmonary disease, asthma, coronary artery disease, cardiomegaly and metabolic syndrome.
The difference in the percentage of comorbidities might be attributed to race and the study population. Because of matching, the difference in baseline characteristics between the uveitis and non-uveitis groups was not statistically significant (Table 1). Therefore, the higher cumulative risk of developing AMI in the uveitis group demonstrated by the Kaplan–Meier curve (Figure 2) is less likely to be attributed to the differences in sex, age, and baseline systemic disease.

**TABLE 4** | Factors for AMI among different uveitis subgroup evaluated by Cox regression with/without Fine & Gray’s competing risk model.

| Uveitis subgroup | Populations | Events | PYs | Rate (per 10^5 PYs) | Adjusted HR | 95% CI | P | Adjusted HR | 95% CI | P |
|------------------|-------------|--------|-----|---------------------|-------------|-------|---|-------------|-------|---|
| Without uveitis  | 4,724       | 415    | 46,864.79 | 885.53           | 1.565       | 1.404 | 1.721 | <0.001*       | 1.633 | 1.480 | 1.818 | <0.001*       |
| With uveitis     | 1,181       | 138    | 10,562.22 | 1,306.54         | 1.597       | 1.435 | 1.789 | <0.001*       | 1.688 | 1.513 | 1.969 | <0.001*       |
| Anterior uveitis | 432         | 52     | 4,292.89  | 1,211.31         | 1.522       | 1.366 | 1.701 | <0.001*       | 1.608 | 1.442 | 1.791 | <0.001*       |
| Posterior segment involvement | 749 | 86 | 6,269.33 | 1,371.76 | 1.595 | 1.436 | 1.848 | <0.001* | 1.673 | 1.510 | 1.930 | <0.001* |

*indicates statistical significance, p < 0.05. PYs, Person-years; Adjusted HR, Adjusted Hazard ratio: adjusted for the variables listed in Table 2; CI, confidence interval; Competing risk: all-cause mortality.
Table 2 shows the variables associated with the development of AMI by Cox regression. As expected, patients with traditional risk factors had a higher risk of developing AMI, a result that is similar to those of previous studies (29, 30). Bremander et al. reported elevated morbidity ratios for hypertension and diabetes in AS patient with ischemic heart disease (29) Chou et al. found that AS patients with hypertension, diabetes, hyperlipidemia and stroke had an aHR of 7.74 for acute coronary syndrome compared to those without these comorbidities (30).

Our report also identified uveitis as an independent risk factor for AMI in patients with AS (Table 3). The reason why uveitis might be a risk factor for developing AMI in patients with AS is not clear. Uveitis may be an indicator of disease activity and systemic inflammation in AS (9, 31, 32). Patients with acute anterior uveitis have higher disease activity and inflammatory cytokines in the serum (9, 33). Since inflammation plays an important role in myocardial infarction (7), uveitis may be associated with cardiovascular disease. A cross-sectional study reported that AS patients with a history of uveitis were more predisposed to cardiovascular disease (34). Another study also reported that children with a longer duration of uveitis had an increased frequency of deteriorated vascular function and carotid artery intima-media thickness (35).

Previous studies have reported that uveitis in spondyloarthritis is associated with HLA-B27, possibly acting through the IL-23/IL-17 axis (36–38). The activation of Th17 cells with Th17-related proinflammatory cytokine induction (IFN-γ, TNF-α, IL-1β, IL-6, IL-23) could result in spondyloarthritis with associated uveitis (39). Several studies have reported an association between Th17-related proinflammatory cytokines and atherosclerosis. A thin fibrous cap in the lesion, which is indicative of unstable plaques prone to rupture and increased atherosclerosis correlate with increased Th17 cells, IL-23-producing vascular muscle, and macrophages in apoE/IL18 double-KO mice (40). Circulating Th17-related cytokines (IL-17, IL-6, IL-23) increased at the onset of acute coronary syndrome (41). The blockade of IL-17A in mice resulted in a significant reduction in atherosclerosis (42, 43). Thus, the IL-23/IL-17 axis is a possible common pathway of origin in uveitis and AMI.

We further divided uveitic patients into the anterior uveitic and posterior segment involvement groups (Table 4). Our study showed that the posterior segment involvement group accounted for the majority (63.4%) of the uveitis cohort. Most studies focused on the association between anterior uveitis and AS. Few studies have discussed the manifestation of posterior segment involvement in patients with AS uveitis. The percentage of posterior segment involvement in HLA-B27 associated uveitis was reported to be 17.4–63%, varying from one study to another (44–47). This difference might be attributed to race, study design, and study year. Our study also showed that the posterior segment involvement group had slightly increased aHR of developing AMI than the anterior uveitis group; however, we did not perform statistical analysis to compare the two groups. Due to the lack of further statistical information, the interpretation of higher aHR in the posterior segment involvement group should be done carefully.

Our study has several strengths. First, the National Health Administration in Taiwan checks charts to ensure patient diagnosis with appropriate treatment. Treatment followed a standardized protocol. The diagnoses were verified. Second, we used a comprehensive, longitudinal, nationwide database spanning 15 years. The NHI system was introduced in Taiwan in 1995, so we could conduct a longitudinal analysis of sequential events. AS or AMI before the index date could be excluded to eliminate bias in the cross-sectional study. Third, the coverage rate in Taiwan was approximately 99%. A large sample provides high statistical power. Finally, the major strength was that the findings after adjusting for confounders were highly consistent, suggesting reliable and convincing results.

However, this study had some limitations. First, the study was based on a retrospective statistical analysis from a database. The NHIRD database does not provide laboratory data, such as the erythrocyte sedimentation rate or C-reactive protein, as advocatory evidence. There were no clinical images such as MRI findings for classification of disease activity and severity of AS patients, which might also show a correlation with uveitis. Second, some traditional factors for AMI, such as diet, smoking, and physical activity, were not in the database. Third, AS-uveitis complex is most frequently observed as anterior uveitis and responds well to topical corticosteroids, whereas posterior uveitis is usually refractory to topical treatments and frequently requires treatments with systemic corticosteroids and anti-TNF agents to reduce recurrences. Cardiovascular events are complications of corticosteroids. In this study, we did not assess the influence of uveitis treatments in developing AMI. Fourth, the rates of all extra-articular manifestation in AS patients might influence the development of AMI, but the database did not provide the information of all extra-articular manifestation. Fifth, the included patients in our study are homogenous ethnic population with specific genetic and disease features as indicated by the prevalence of comorbidities, type of uveitis and probably age at diagnosis. Hence, the results of our study must be confirmed in other ethnic populations. Sixth, this study had a retrospective design. We could only elucidate a possible mechanism based on this association. The causality needs further prospective studies with detailed information to confirm our conclusion.

CONCLUSION

Uveitis is a risk factor for developing AMI in patients with AS. Doctors should be aware of the potential cardiovascular risk in patients with AS uveitis, especially simultaneously with other traditional risk factors for AMI. Closely monitoring the cardiovascular system in these patients is important for all physicians to detect AMI earlier and manage it promptly. Further prospective studies are needed to elucidate the underlying mechanism between uveitis and AMI in patients with AS.
DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/Supplementary Material. Further inquiries can be directed to the corresponding author.

ETHICS STATEMENT

The studies involving human participants were reviewed and approved by The institutional review board of the Tri-Service General Hospital (TSGH IRB No. B-110-35). Written informed consent for participation was not required for this study in accordance with the national legislation and the institutional requirements.

AUTHOR CONTRIBUTIONS

Conceptualization: Y-FL, W-CC, and C-LC. Methodology: W-CC and C-HC. Software: W-CC, C-AS, and C-HC. Validation: T-YL, W-CC, and C-HC. Formal analysis: W-CC, C-AS, and C-HC. Investigation: T-YL, Y-HC, and J-TC. Resources: W-CC, Y-HC, J-TC, and C-LC. Data curation: C-AS, and C-HC. Investigation: T-YL, Y-HC, and J-TC. Validation: T-YL, W-CC, and C-HC. Formal analysis: W-CC, W-CC and C-HC. Software: W-CC, C-AS, and C-HC.

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fimmu.2021.811664/full#supplementary-material

REFERENCES

1. Feldtkeller E, Khan MA, van der Heijde D, van der Linden S, Braun J. Age at Disease Onset and Diagnosis Delay in HLA-B27 Negative vs. Positive Patients With Ankylosing Spondylitis. Rheumatol Int (2003) 23(2):61–6. doi: 10.1007/s00296-002-0237-4
2. Luo F, Zhao Z, Zhang J, Leng J. Comparison of HLA-B*27 Subtypes Between Chinese Patients With Ankylosing Spondylitis and Non-Ankylosing Spondylitis Carriers. J Int Med Res (2019) 47(7):3171–8. doi: 10.1177/0300060519853929
3. Prati C, Demougout C, Guillo X, Sondag M, Verhoeven F, Wending D. Vascular Involvement in Axial Spondyloarthropathies. Joint Bone Spine (2019) 86(2):159–63. doi: 10.1016/j.jbspin.2018.05.003
4. Mathieu S, Pereira B, Soubrier M. Cardiovascular Events in Ankylosing Spondylitis: An Updated Meta-Analysis. Semin Arthritis Rheumat (2015) 44(5):551–5. doi: 10.1016/j.semarthrit.2014.10.007
5. Eriksson JK, Jacobsen L, Bengtsson K, Askling J. Is Ankylosing Spondylitis a Risk Factor for Cardiovascular Disease, and How Do These Risks Compare With Those in Rheumatoid Arthritis? Ann Rheum Dis (2017) 76(2):364–70. doi: 10.1136/annrheumdis-2016-209315
6. Huang YP, Wang YH, Pan SL. Increased Risk of Ischemic Heart Disease in Young Patients With Newly Diagnosed Ankylosing Spondylitis–A Population-Based Longitudinal Follow-Up Study. PloS One (2013) 8(5): e64155. doi: 10.1371/journal.pone.0064155
7. Hansson GK. Inflammation, Atherosclerosis, and Coronary Artery Disease. N Engl J Med (2005) 352(16):1685–95. doi: 10.1056/NEJMra043430
8. Zeboulon N, Dougados M, Gossec L. Prevalence and Characteristics of Uveitis in the Spondyloarthropathies: A Systematic Literature Review. Ann Rheumat Dis (2008) 67(7):955–9. doi: 10.1136/ard.2007.075774
9. Jawad S, Liu B, Agron E, Nussenblatt RB, Sen HN. Elevated Serum Levels of Interleukin-17A in Uveitis Patients. Ocular Immunol Inflamm (2013) 21(6):434–9. doi: 10.3109/09273949.2013.815786
10. Curnow SJ, Falciani F, Durrani OM, Cheung CM, Ross EJ, Wloka K, et al. Multiplex Beat Immunoassay Analysis of Aqueous Humor Reveals Distinct Cytokine Profiles in Uveitis. Invest Ophthalmol Vis Sci (2005) 46(11):4251–9. doi: 10.1167/iovs.05-0444
11. Carreno E, Portero A, Herreras JM, Garcia-Vazquez C, Whitcup SM, Stern ME, et al. Cytokine and Chemokine Tear Levels in Patients With Uveitis. Acta Ophthalmol (2017) 95(5):e405–14. doi: 10.1111/aos.13292
12. Dean LE, Jones GT, MacDonald AG, Downham C, Sturrock RD, Macfarlane GJ. Global Prevalence of Ankylosing Spondylitis. Rheumatol (Ox Engl) (2014) 53(4):650–7. doi: 10.1093/rheumatology/ket387
13. Osten F, Akar S, Birlik M, Sari I, Khan MA, Gurler O, et al. Prevalence of Ankylosing Spondylitis and Related Spondyloarthritides in an Urban Area of Izmir, Turkey. J Rheumatol (2008) 35(2):305–9.
14. Chou CT, Pei L, Chang DM, Lee CF, Schumacher HR, Liang MH. Prevalence of Rheumatic Diseases in Taiwan: A Population Study of Urban, Suburban, Rural Differences. J Rheumatol (1994) 21(2):302–6.
15. Joshi VL, Chopra A. Is There an Urban-Rural Divide? Population Surveys of Ankylosing Spondylitis and Related Spondyloarthritides in an Urban Area of Izmir, Turkey. J Rheumatol (2008) 35(2):305–9.
16. Alvarez-Nemegyev J, Peláez-Ballestas I, Sanin LH, Cardiel MH, Ramirez-Angulo A, Goyochoa-Robles MV. Prevalence of Musculoskeletal Pain and Rheumatic Diseases in the Southeastern Region of Mexico. A COPCORD-Based Community Survey. J Rheumatol Suppl (2011) 86:21–5. doi: 10.3899/jrheum.100954
17. De Angelis R, Salaﬁ F, Grassi W. Prevalence of Spondyloarthropathies in an Italian Population Sample: A Regional Community-Based Study. Scand J Rheumatol (2007) 36(1):14–21. doi: 10.1080/03009740600904243
18. Davatchi F, Jamshidi AR, Banishahemi AT, Gholami J, Forouzanfar MH, Akhlaghi M, et al. WHO-ILAR COPCORD Study (Stage 1, Urban Study) in Iran. J Rheumatol (2008) 35(7):1384.
19. Trontzas P, Andraniakos A, Miyakis S, Pantelidou K, Vafiadou E, Garantziotou V, et al. Seronegative Spondyloarthropathies in Greece: A Population-Based Study of Prevalence, Clinical Pattern, and Management. The ESORDIG Study. Clin Rheumatol (2005) 24(6):583–9. doi: 10.1007/s10067-005-1106-9

20. Cakır N, Pamuk ÖN, Derviş E, Imeryüz N, Ulu H, Benian Ö, et al. The Prevalences of Some Rheumatic Diseases in Western Turkey: Hava Study. Rheumatol Int (2012) 32(4):895–908. doi: 10.1007/s00296-010-1699-4

21. Stolwijk C, van Tubergen A, Castillo-Ortiz JD, Boonen A. Prevalence of Extra-Articular Manifestations in Patients With Ankylosing Spondylitis: A Systematic Review and Meta-Analysis. Ann Rheum Dis (2015) 74(1):65–75. doi: 10.1136/annrheumdis-2013-205382

22. Hsu YR, Huang JC, Tao Y, Kaburaki T, Lee CS, Lin TC, et al. Noninfectious Uveitis in the Asia-Pacific Region. Eye (Lond Engl) (2019) 33(1):66–77. doi: 10.1038/s41333-018-0223-z

23. Chen SC, Chuang CT, Chu MY, Sheu SJ. Patterns and Etiologies of Uveitis at a Tertiary Referral Center in Taiwan. Ocular Immunol Inflamm (2017) 25(sup1):S1–8. doi: 10.1080/09273948.2016.1189577

24. Abaño JM, Galvante PR, Siopongco P, Dans K, Lopez J. Review of Atopic Disorders in Ankylosing Spondylitis and Rheumatoid Arthritis. Clin Immunol (2018) 200:102679. doi: 10.1016/j.clim.2018.04.002

25. Haroon NN, Paterson JM, Li P, Inman RD, Haroon N. Patients With Ankylosing Spondylitis Have Increased Cardiovascular and Cerebrovascular Mortality: A Population-Based Study. Ann Intern Med (2015) 163(6):409–16. doi: 10.7326/M14-2470

26. Sharif K, Watad A, Tiosano S, Yavne Y, Blokh Kerpel A, Comaneshter D, et al. The Link Between COPD and Ankylosing Spondylitis: A Population Based Study. Eur J Intern Med (2018) 53:62–5. doi: 10.1016/j.ejim.2018.04.002

27. Singh DK, Magrey MN. Racial Differences in Clinical Features and Comorbidities in Ankylosing Spondylitis in the United States. J Rheumatol (2020) 47(6):835–8. doi: 10.3899/jrheum.181019

28. Rudwaleit M, Andermann B, Alten R, Sörensen H, Listing J, Zink A, et al. Blockade of Interleukin-17A Results in Reduced Atherosclerosis in Apolipoprotein E-Deficient Mice. Circulation (2010) 121(15):1746–55. doi: 10.1161/CIRCULATIONAHA.109.248866

29. Cheng X, Yu X, Ding YJ, Fu QQ, Xie JJ, Tang TT, et al. The Th17/Treg Imbalance in Patients With Acute Coronary Syndrome. Clin Immunol (Orlando Fla) (2008) 127(1):89–97. doi: 10.1016/j.clim.2008.01.009

30. Smith E, Prasad KM, Butcher M, Dobrian A, Kolls JK, Ley K, et al. Blockade of Interleukin-17A Attenuates Atherosclerotic Lesion Development in apoE-Deficient Mice. J Immunol (Baltimore Md: 1950) (2009) 183(12):8167–75. doi: 10.4049/jimmunol.0901126

31. Bayen H, Bayen MC, De Curzon HP, Espinasse-Berrod MA, Mandiereux N, Furia M, et al. Involvement of the Posterior Eye Segment in HLA B27(+)-Iridocyclitis. Incidence. Value of Surgical Treatment. J Fransais D Ophtalmol (1988) 11(8):561–6

32. Rodriguez A, Akova YA, Pedroza-Seres M, Foster CS. Posterior Segment Ocular Manifestations in Patients With HLA-B27-Associated Uveitis. Ophthalmology (1994) 101(7):1267–74. doi: 10.1016/S0161-6420(94)31179-1

33. Chou CH, Lin MC, Peng CL, Wu YC, Sung FC, Kao CH, et al. A Nationwide Population-Based Retrospective Cohort Study: Increased Risk of Acute Coronary Syndrome in Patients With Ankylosing Spondylitis. Scand J Rheumatol (2014) 43(2):132–6. doi: 10.3109/03009742.2013.822097

34. Chen CH, Lin KC, Chen HA, Liao HT, Liang TH, Wang HP, et al. Association of Acute Anterior Uveitis With Disease Activity, Functional Ability and Physical Mobility in Patients With Ankylosing Spondylitis: A Cross-Sectional Study of Chinese Patients in Taiwan. Clin Rheumatol (2007) 26(6):953–7. doi: 10.1007/s10067-006-0403-2

35. Robertson LP, Davis MI. A Longitudinal Study of Disease Activity and Functional Status in a Hospital Cohort of Patients With Ankylosing Spondylitis. Rheumatol (Oxf) (2004) 43(12):1565–8. doi: 10.1093/rheumatology/keh386

36. Martin TM, Rosenbaum JT. An Update on the Genetics of HLA B27-Associated Acute Anterior Uveitis. Ocular Immunol Inflamm (2011) 19(2):108–14. doi: 10.3109/09273948.2011.593032

37. Delay ML, Turner MJ, Klenk EI, Smith JA, Sowders DP, Colbert RA. HLA-B27 Misfolding and the Unfolded Protein Response: A Novel Interleukin-23 Production and Activation. Arthritis Rheumatol (2009) 60(9):2633–43. doi: 10.1002/art.24763

38. Zhong Z, Su G, Kijistra A, Yang P. Activation of the Interleukin-23/Interleukin-17 Signalling Pathway in Autoinflammatory and Autoimmune Uveitis. Prog Retin Eye Res (2021) 80:100866. doi: 10.1016/j.preteyeres.2020.100866

39. Feldkamp D, Yates W, Amjadi S, McCluskey P. HLA-B27 Anterior Uveitis: Immunology and Immunopathology. Ocular Immunol Inflamm (2016) 24(4):450–9. doi: 10.3109/09273948.2016.1158283

40. Pejnovic N, Vratimos A, Lee SH, Popadic D, Takeda K, Akira S, et al. Increased Atherosclerotic Lesions and Th17 Deficiency in Apolipoprotein E Knockout Mice. Mol Immunol (2009) 47(1):37–45. doi: 10.1016/j.molimm.2008.12.032