Daily high-dose aspirin does not lower APRI in the Aspirin-Myocardial Infarction Study

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Abstract

Antiplatelet agents reduce liver fibrosis by inhibiting platelet activation and platelet-derived growth factor production. Previous cross-sectional epidemiological studies suggest that the use of aspirin is related to reduced liver fibrosis. The Aspirin-Myocardial Infarction Study (AMIS) aims to examine this relationship in a multicenter, randomized, double-blind and placebo-controlled trial. The existing clinical trial of aspirin was conducted to study the benefit of one gram aspirin daily among 4524 individuals who had experienced at least one documented myocardial infarction. The aspartate aminotransferase (AST)-to-Platelet Ratio Index (APRI) was calculated at baseline and annually from the platelet count and AST levels. Participants in the AMIS trial had a mean baseline APRI of 0.34±0.36, and only 1% individuals had APRI scores higher than 1.0, a common cutoff for cirrhosis. The daily use of aspirin was associated with an increase, rather than a reduction of APRI, by 0.007 per year (95% CI 0.002−0.015, P=0.12). The use of aspirin did not significantly affect platelet counts. In a sensitivity analysis of individuals with probable significant fibrosis at baseline (APRI≥0.7), the aspirin group had a sustained reduction in APRI over time, although this change was not significant compared to that in the placebo group. In the AMIS trial, the daily use of high-dose aspirin did not significantly affect APRI, a surrogate index of liver fibrosis. This study highlights the need for de novo clinical trials to investigate the potential benefit of antiplatelet agents on liver fibrosis.

Keywords: liver fibrosis, aspirin, APRI, antiplatelet, AMIS

Introduction

Liver fibrosis is a common pathway for most forms of chronic liver disease that leads to cirrhosis and liver failure. In addition to its well-established function in thrombus formation, platelets play a crucial role in the regulation of inflammation and tissue repair[1]. In the liver, platelet-derived growth factor-beta (PDGF-β) is a key profibrogenic cytokine that promotes the activation of quiescent stellate cells into myofibroblasts[2–3]. In addition, activated platelets release tumor growth factor-beta and chemokine (C-X-C motif) ligand 4. Together with PDGF-β, these cytokines can promote fibrogenesis in liver diseases through the activation of quiescent hepatic stellate cells[4–9]. As a result, antiplatelet therapy, antibody-
mediated platelet depletion, and aspirin protect against liver fibrosis in murine models of liver fibrosis[4]. Similarly, aspirin, ticlopidine, and cilostazol, three antiplatelet agents, attenuate liver steatosis, inflammation and fibrosis in diet-induced rat models of nonalcoholic fatty liver disease (NAFLD)[6]. Data from the National Health and Nutrition Examination Survey (NHANES), a nationally representative cross-sectional study, suggested that individuals using aspirin, especially those with chronic liver diseases, had lower indices of liver fibrosis compared to those with no aspirin use[7]. This putative anti-fibrotic association was not seen in those with the use of ibuprofen[7]. A recent study by Schwarzkopf et al demonstrated an inverse association between the use of antiplatelet agents (aspirin and/or clopidogrel) and liver fibrosis measured by transient elastography among patients with cardiovascular diseases[8]. In their study, the circulating level of PDGF-β was associated with platelet counts but not with the use of antiplatelet agents. Furthermore, a prospective study by Simon and colleagues showed that among individuals with NAFLD and low-grade fibrosis (stage 0–2), the daily use of aspirin is associated with a significantly lower risk of incidental development of advanced liver fibrosis (stage 3–4) measured by noninvasive fibrosis indices[9]. The putative anti-fibrotic effect of antiplatelet agents, such as aspirin, has not yet been studied in clinical trials. Here, we examine the prospective impact of daily aspirin use on liver fibrosis measured by the aspartate aminotransferase (AST)-to-Platelet Ratio Index (APRI) in the Aspirin-Mycardial Infarction Study (AMIS).

Materials and methods

AMIS was a multicenter, randomized, double-blind, placebo-controlled trial, sponsored by National Heart, Lung and Blood Institute and conducted to study the impact of aspirin on mortality among individuals with a documented history of myocardial infarction[10]. A total of 4,524 men and women aged between 30 and 68 in the United States were randomized to receive either one gram of aspirin or placebo daily for three years. Platelet count and AST were measured at baseline and annually at follow-up visits. Primary data were provided by NHLBI. APRI, a validated fibrosis index, was measured as previously described[11]. We analyzed the association between aspirin use and changes in APRI over time using generalized estimating equations and an unstructured correlation matrix, in which the impact of aspirin was calculated as the time-by-treatment interaction between the year of follow-up (0–3) and aspirin assignment. A sensitivity analysis was conducted to examine the impact of aspirin on subjects likely to have significant liver fibrosis defined by APRI≥0.7 at baseline. All statistical analyses were performed using Stata version 13 (College Station, USA).

Results

The baseline characteristics of patients in the AMIS trial are summarized in Table 1. Most study participants were Caucasian (92%) and male (~89%) with a mean age of 55 years, body-mass index of 26, and a prevalence of diabetes of 2.6%. The mean APRI values at baseline were 0.34±0.46 in the placebo and 0.33±0.22 in the aspirin groups. Only 2.5% of individuals (n=113) had baseline APRI higher than 0.7.

Over three years, mean APRI increased in both aspirin and placebo groups compared to baseline. Assignment to aspirin tended to be associated with a greater increase in the rate of change in APRI over time than placebo (difference in the rate of change 0.007 per year; 95% CI 0.002–0.015, P=0.12) (Fig. 1A). There was no significant difference in the rate of

| Table 1 Baseline characteristics of study participants |
|-----------------------------------------------|
| Characteristic | Placebo (n=2,257) | Aspirin (n=2,267) |
|----------------|------------------|------------------|
| Age (years)   | 54.9±7.9         | 54.8±8.1         |
| Male          | 89.4%            | 88.4%            |
| Ethnicity     |                  |                  |
| White         | 91.5%            | 91.7%            |
| Black         | 5.9%             | 6.3%             |
| Hispanics     | 2.6%             | 2.0%             |
| BMI           | 26.0±3.5         | 25.9±3.5         |
| Diabetes      | 2.8%             | 2.5%             |
| Smoking       |                  |                  |
| Never         | 27.2%            | 27.5%            |
| Former        | 5.4%             | 5.3%             |
| Current       | 19.2%            | 19.7%            |
| Alcohol consumption |        |                  |
| Daily drink (>1.5 ounces) | 15.4% | 15.9% |
| Laboratory results |                |                  |
| Platelet count (<10^11/L) | 253±70 | 257±70 |
| AST (IU/L)    | 32±34            | 31±15            |
| ALP (IU/L)    | 62±22            | 64±23            |
| Bilirubin (mg/dL) | 0.7±0.3 | 0.7±0.3 |
| APRI          | 0.34±0.46        | 0.33±0.22        |
changes in platelet counts between the aspirin and placebo groups over time ($P=0.4$), but a trend toward 0.52 IU/L higher AST annually in the aspirin group compared to the placebo group (95% CI 0.05 –1.09, $P=0.07$) was noted. Among individuals with elevated APRI at baseline (mean APRI of 1.1±0.5 for the aspirin and 1.0±0.4 for the placebo group), APRI values tended to decrease more consistently over time among subjects who received aspirin than among those who received placebo (Fig. 1B), but the differences in changes of APRI over time were not significant between the two groups ($P=0.7$).

Discussion

Current strategies of treating fibrosis in chronic liver diseases anchor on the treatment of underlying causes to prevent disease progression. There is an urgent need for therapeutic interventions that can specifically target fibrogenesis and potentially reverse fibrosis. Accumulating evidence suggests that antiplatelet agents, such as aspirin, may have anti-fibrotic properties and could be a cost-effective therapeutic option.

Randomized clinical trials testing this therapeutic indication of aspirin are yet to be conducted. Very few are antiplatelet clinical trials with liver-related measurements. AMIS allowed us to leverage the annual measurement of the platelet count and AST to calculate APRI, a validated index of liver fibrosis. In this study, daily use of 1 000 mg of aspirin did not change the three-year trajectory of APRI compared with the placebo. Although APRI is an imperfect index for liver fibrosis, these results from a large randomized trial suggest an absence of sizable benefit from daily aspirin in the primary prophylaxis for liver fibrosis among individuals without pre-existing liver diseases. On the contrary, there was a trend toward higher APRI among aspirin users over time, albeit small and unlikely to be clinically significant. AMIS used 1 000 mg of aspirin in the treatment arm, a practice common at the time of the clinical trial, but significantly higher than current standards. As the inhibition of aspirin on platelets is irreversible, a higher dose is not likely to render additional benefits but could result in significant side effects. Of note, idiosyncratic hepatotoxicity due to aspirin can occur, especially at the high dose used in AMIS. Therefore, a liver protective effect of aspirin could manifest at lower doses of aspirin at 75 –100 mg as used for cardiovascular prevention. The dose-effectiveness of aspirin for liver fibrosis has not been formally studied.

The AMIS trial does not directly address the impact of aspirin among individuals with chronic liver disease, as few participants had significant pre-existing fibrosis by APRI. The negative result does not negate the preclinical observations of platelet involvement in the regulation of liver inflammation and fibrosis. In one retrospective study, low-dose daily aspirin was associated with a lower risk of liver fibrosis progression among patients with HCV recurrence after liver transplantation[12]. Others have noted that an association between aspirin use and reduced risks of hepatocellular carcinoma in a mouse model of chronic hepatitis and human studies[13]. In AMIS, there was a numerical tendency toward lower APRI values over time with aspirin use in the subgroup of subjects with high APRI values, but this result was far from significant and limited by regression to the mean among those with initially high APRI values.

As a secondary analysis, this study has major limitations that include a patient population not representative of patients with chronic liver diseases, an imperfect proxy fibrosis measurement, a dose of aspirin higher than the current standard and therefore unable to minimize its adverse effects, and a moderate duration of follow-up. A negative result here
highlights the need for de novo clinical trials to investigate the potential benefit of antiplatelet agents for chronic liver diseases. Nonetheless, observations here can provide useful insights for future trial design. Foremost, the putative antifibrotic benefit of antiplatelets such as aspirin is likely minimal among individuals without chronic liver diseases. This was also seen in NHANES III where the negative association between aspirin use and fibrosis indices was significantly larger among individuals with chronic liver diseases than in those with no liver diseases[7]. Hence, a primary prophylaxis trial, such as the design in the Physician Health Study, may not be appropriate. The modality of fibrosis measurement and the duration of follow-up are also important considerations that need to be weighed against the costs of the study.

Overall, we report a negative association between the daily use of high-dose aspirin and changes in APRI over three years among patients with a prior history of myocardial infarction. Despite encouraging preclinical and cross-sectional studies, randomized clinical trials are necessary to evaluate the potential therapeutic value of antiplatelet agents in treating liver fibrosis in chronic liver diseases.

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