Risk of atrial fibrillation in athletes: a systematic review and meta-analysis

William Newman, Gemma Parry-Williams, Jonathan Wiles, Jamie Edwards, Sabina Hulbert, Konstantina Kipourou, Michael Papadakis, Rajan Sharma, Jamie O’Driscoll

ABSTRACT

Objective A systematic review, meta-analysis and meta-regression were performed on selected studies to investigate the incidence of atrial fibrillation (AF) among athletes compared with non-athlete controls.

Design Meta-analysis with heterogeneity analysis and subsequent meta-regression to model covariates were performed. The mode of exercise (endurance and mixed sports) and age were the a priori determined covariates.

Data sources PubMed, MEDLINE, SPORTDiscus and the Cochrane library were searched.

Eligibility criteria Research articles published after 1990 and before 2 December 2020 were included if they reported the number of AF cases in athletes with non-athlete physically active or inactive control groups, were case–control or cohort studies and if data allowed calculation of OR.

Results The risk of developing AF was significantly higher in athletes than in non-athlete controls (OR: 2.46; 95% CI 1.73 to 3.51; p<0.001, I²=4.97). Mode of exercise and risk of AF were moderately correlated (B=0.1259, p=0.0193), with mixed sport conferring a greater risk of AF than endurance sport (B=−0.02476, p=0.0204). Younger (<55 years) athletes were significantly more likely to develop AF compared with older (≥55 years) athletes (B=−0.02293, p<0.001).

Conclusion Athletes have a significantly greater likelihood of developing AF compared with non-athlete controls, with those participating in mixed sport and younger athletes at the greatest risk. Future studies of AF prevalence in athletes according to specific exercise dose parameters, including training and competition history, may aid further in delineating those at risk.

INTRODUCTION

Physical activity and exercise training reduce metabolic disease risk, improve cardiovascular health and mental well-being and are associated with healthy ageing. Studies also suggest that exercise can reduce the burden of arrhythmias, such as atrial fibrillation (AF), which is the most common cardiac arrhythmia worldwide. Indeed, the prevalence of AF in the general population is as high as 3.3% and is associated with a 1.5-fold increased mortality risk, most commonly secondary to cerebrovascular events. AF is also the most prevalent arrhythmia in athletic populations. It has been reported that male endurance athletes performing high-intensity and high-volume exercise training are at an increased risk, with studies suggesting a U-shaped curve, with lack of physical activity and long-term vigorous exercise both increasing the risk of developing AF. The mechanisms by which exercise training increases the risk of AF are complex and speculative, but may include atrial dilation, adrenergic activation, vagal tone, chronic inflammation, pulmonary foci and interstitial fibrosis, occurring as a result of excessive strain through augmented cardiac output and atrial stretch.

Two prior meta-analyses investigating the possible increased risk of AF in athletes have demonstrated equivocal findings, with ORs of 1.64 and 5.29. Interestingly, Ayinde et al identified age as a significant determinant of AF risk, with younger athletes (below 40 years old) at greater relative risk than athletes >34 years of age. However, these meta-analyses excluded studies of athletes with hypertension or diabetes, which potentially restricts the availability of larger pooled study data for analysis. As such, this meta-analysis aims to establish the risk of AF in athletes and further elucidate the influence of sport type and cardiovascular disease (CVD) risk factors on the risk of developing AF. As the prevalence is relatively low, this paper aimed to perform a meta-analysis and pool multiple study data to draw a more conclusive overall prevalence among athletes.

METHODS

Search strategy

This systematic review and meta-analysis was fully compliant with and performed according to the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) guidelines, including use of the described PICOS method for the systematic review process. Figure 1 details the PRISMA systematic review flowchart.

PubMed, MEDLINE, Science Direct, the Cochrane library and SPORTDiscus online databases were searched for studies reporting AF or atrial flutter in athletes in journal articles published before 2 December 2020. Studies written in English and published from 1990 onwards were accepted. Key Boolean search terms were utilised to identify relevant studies and included; ‘atrial fibrillation’ OR ‘atrial flutter’ AND ‘athlete’, ‘sport’, ‘endurance’ AND ‘exercise’. Where possible, corresponding study authors were contacted to ascertain whether non-published data was available or in the pre-print stage, and studies found through the systematic protocol were screened for exterior citations and their respective reference lists searched for eligible studies.
Review

Study eligibility

Screening was carried out following the exclusion of duplicate articles. Studies were initially excluded by title and then by abstract. Studies retained for the next step of evaluation were screened and included if: they reported the number of AF cases in athletes with non-athlete control groups, were case–control or cohort study design and the presentation of data was in the form of ORs, or at least data whereby OR and CIs could be precisely calculated.

Studies were excluded in the next evaluation step if they did not include a control group and participants older than 18 years who had performed regular exercise training for at least a 2-year period prior to screening. For the purpose of this meta-analysis, ‘athletes’ were defined according to the European Society of Cardiology classification.17 Studies of athlete’s participating in sports other than the following were excluded: cycling, running, swimming, Nordic skiing, orienteering, rowing (endurance sports) and mixed sports (football, netball, rugby, etc) as defined by Pelliccia et al.17 Studies where participants had concurrent diseases were excluded except for hypertension and diabetes, to exclude other confounding causes of AF but to include conditions common among the general population.18 Studies with these two concurrent diseases were classified and grouped for further subgroup analysis to determine if these diseases moderate the degree of AF prevalence and effect. Furthermore, studies reporting atrial flutter were included due to it being a precursor to AF.

For this meta-analysis, we included studies with non-exercising or recreationally exercising controls versus athlete populations to ascertain whether an optimal exercise duration, intensity or mode exists before AF manifests versus the general population. Non-athletic leisurely activity and physical activity participants were defined as physically active individuals without training regimes, athletic ambitions or participation in regular competition. In the literature, this has been regarded as ‘leisure time physical activity’. This type of physical activity is generally at or below national physical activity guidelines, but not near the volumes or intensities seen through purposed exercise training undertaken by athletes.

Data extraction, coding and study quality

Our initial search identified 3885 citations. After all exclusions were applied, 13 studies were included and proceeded through to the data extraction and coding step (figure 1). The 13 studies included in the meta-analysis (table 1) were coded and

Table 1 Characteristics of studies included

<table>
<thead>
<tr>
<th>Author</th>
<th>Year</th>
<th>Country</th>
<th>Sample size</th>
<th>Average age (years)</th>
<th>Sport type</th>
<th>Study design</th>
<th>Risk of bias*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aizer et al</td>
<td>2009</td>
<td>America</td>
<td>8448</td>
<td>52</td>
<td>Mixed and Endurance†</td>
<td>Cohort</td>
<td>2</td>
</tr>
<tr>
<td>Andersen et al</td>
<td>2013</td>
<td>Sweden</td>
<td>52 755</td>
<td>57</td>
<td>Nordic Skiing</td>
<td>Cohort</td>
<td>2</td>
</tr>
<tr>
<td>Baldesberger et al</td>
<td>2008</td>
<td>Switzerland</td>
<td>124</td>
<td>67</td>
<td>Cycling</td>
<td>Cohort</td>
<td>2</td>
</tr>
<tr>
<td>Calvo et al</td>
<td>2016</td>
<td>Spain</td>
<td>172</td>
<td>46</td>
<td>Mixed and Endurance†</td>
<td>Case–control</td>
<td>1</td>
</tr>
<tr>
<td>Claessen et al</td>
<td>2011</td>
<td>Belgium</td>
<td>156</td>
<td>52</td>
<td>Mixed and Endurance§</td>
<td>Case–control</td>
<td>2</td>
</tr>
<tr>
<td>Eloisa et al</td>
<td>2006</td>
<td>Spain</td>
<td>160</td>
<td>43</td>
<td>Mixed and Endurance§</td>
<td>Case–control</td>
<td>1</td>
</tr>
<tr>
<td>Karjalainen et al</td>
<td>1998</td>
<td>Finland</td>
<td>440</td>
<td>46</td>
<td>Orienteering</td>
<td>Cohort</td>
<td>1</td>
</tr>
<tr>
<td>Molina et al</td>
<td>2008</td>
<td>Spain</td>
<td>473</td>
<td>39</td>
<td>Running</td>
<td>Cohort</td>
<td>1</td>
</tr>
<tr>
<td>Mont et al</td>
<td>2002</td>
<td>Spain</td>
<td>216</td>
<td>44</td>
<td>Mixed and Endurance**</td>
<td>Case–control</td>
<td>2</td>
</tr>
<tr>
<td>Myrstad et al</td>
<td>2014</td>
<td>Norway</td>
<td>2376</td>
<td>69</td>
<td>Nordic Skiing</td>
<td>Cohort</td>
<td>2</td>
</tr>
<tr>
<td>Myrstad et al</td>
<td>2016</td>
<td>Norway</td>
<td>4952</td>
<td>69</td>
<td>Nordic Skiing</td>
<td>Cohort</td>
<td>2</td>
</tr>
<tr>
<td>Scheiner et al</td>
<td>2016</td>
<td>America</td>
<td>149</td>
<td>72</td>
<td>Swimming</td>
<td>Case–control</td>
<td>3</td>
</tr>
<tr>
<td>Van Bauren et al</td>
<td>2012</td>
<td>Germany</td>
<td>57</td>
<td>57</td>
<td>Handball</td>
<td>Case–control</td>
<td>2</td>
</tr>
</tbody>
</table>

References are available in the supplemental file.

* Risk of bias: 1=low, 2=moderately low, 3=moderately high, 4=high. See online supplemental file 1 for full references.
†Cycling, racquet sports and swimming.
‡Cycling, racquet sports and swimming.
§Basketball, cycling, diving, handball, horse riding, running, soccer, tennis, triathlon.
¶Unknown sports.
**Athletics, cycling, running, soccer, swimming, weight lifting.
respective data was extracted including systematic processing of study details, country of registration, year of publication, journal impact statistics and full patient anthropometric and clinical variables, where available. Study data was extracted and a risk of bias analysis performed, which were systematically screened by a second reviewer. When disputes were detected in coding or quality analyses, the reviewers met to discuss any conflicts, which were then resolved and evaluated further.

AF OR, relative risks and raw data were extracted, and OR was calculated according to the presence of sample size, control and athlete AF diagnosis data. The adjusted status and variables adjusted for in each study are provided in the (online supplementary table S1). When insufficient data was detected, best efforts were made to contact the relevant authors of the study or to seek online supplemental data. If this data was not available, these studies were then excluded from the rest of the analyses.

Qualitative risk of bias assessment was carried out utilising the tool from CLARITY at McMaster University (Evidence Partners, 2019) to ascertain the risk of bias in the cohort and case-control studies independently. For each of the outcomes, a number was assigned to the qualitative descriptor outcomes. The questions included queries surrounding the sampling of the population, composition and extraction of clinical data, and other factors surrounding blinding of participants and researchers. For case-control studies, there were five questions, whereas eight questions were asked for cohort studies. The response to the questions was completed using Likert integers; for example, in the question ‘can we be confident in the assessment of the exposure’, 1=definitely yes (low risk of bias) and 5=definitely no (high risk of bias). The overall results of the study quality assessment composed of a single numerical value resulting from an average of the individual question outcomes. Further quantitative study quality metrics not addressed by qualitative means were computed alongside the meta-analysis utilising funnel plot analyses and asymmetry testing with tests of heterogeneity.

Data analysis and research synthesis

As aforementioned, raw data was extracted from studies and transformed into OR with 95% CIs. Comprehensive Meta-Analysis (Comprehensive Meta-Analysis V.3, Biostat, Englewood, New Jersey, USA) was used to assess the reported cases of AF in athletes versus non-athletes. A random effects pooled analysis was performed on the 13 final studies due to the inter-study variability with case-control and cohort studies, and the assessment of AF risk in all athletes from a population, allowing conclusions of a more general level of inference. Furthermore, random effects models are typically selected when inter-study variability is confirmed through significant heterogeneity. The results of the pooled analysis were considered significant when a p value of <0.05 was present and the test Z statistic was >2. Meta-analyses were also performed independently on the following conditions; age, comorbid CVD risk factor inclusion, study quality and study design, to investigate pooled effects.

Studies were inputted systematically, allowing the individual weightings of each study to be visible and the weighted effect on the overall effect size of the pooled analysis discerned for each study (online supplementary table S2). Heterogeneity was tested alongside the pooled analysis to quantitatively assess the study quality and risk of bias using the $\chi^2$ technique and reported as the I$^2$ statistic. If the I$^2$ statistic was $>40\%$, it was considered significant. Once past this threshold, post hoc tests such as Egger’s test (1997) were systematically planned to qualitatively assess the presence of funnel plot asymmetry, suggesting publication bias.

A meta regression was run to ascertain if any effect moderator variables influenced the risk of developing AF in athletes versus non-athletes. The planned moderators to be assessed independently were presence of diabetes, presence of hypertension, number of cases of high cholesterol, cases of cigarette smoking, body mass index, mode of exercise, age, sport type and study design.

RESULTS

Study and participant characteristics

The meta-analysis included 13 studies (7 cohort and 6 case-control) with a pooled sample size of 70,478 participants, which included 63,662 controls and 6,816 athletes (online supplemental file). Five studies included athletes with the CVD risk factors diabetes and/or hypertension. Risk of bias scores and sport types are presented in Table 1. Only two studies provided data on volume or average duration of exercise and both defined this by lifetime exercise hours and could therefore not be used to quantify weekly exposure.

Primary analysis

The initial random effects pooled analysis demonstrated that the risk of AF was significantly higher in athletes than non-athletes (OR: 2.46; 95% CI 1.73 to 3.51, p<0.001, Z=4.97) (figure 2). There was significant heterogeneity among the studies (p<0.001, I$^2$=84%). The post hoc Egger’s test (1997) was statistically significant (p<0.001), suggesting publication bias.

Subgroup analysis

When dichotomising studies into those with and without CVD risk factors (type 2 diabetes mellitus and hypertension), there was no significant difference in the relative risk of AF in athletes with CVD risk factors versus non-athletes with CVD risk factors (OR: 1.51; 95% CI 0.90 to 2.54, p=0.116, Z=1.572). However, in athletes and non-athletes without CVD risk factors, athletes had a significantly greater relative risk of AF (OR: 3.60; 95% CI 2.28 to 5.88, p<0.001, Z=3.355). Younger athletes (OR: 3.60; 95% CI 2.09 to 6.29, p<0.001, Z=4.603) had a significantly higher relative risk of AF than older athletes (OR: 1.76; 95% CI 0.97 to 3.21, p=0.065, Z=1.845) (figure 3) and good quality studies (OR: 2.32; 95% CI 1.62 to 3.22, p<0.001, Z=4.610) reported lower overall OR for AF in athletes than poor quality studies (OR: 4.80; 95% CI 1.22 to 18.88, p=0.025, Z=2.244) (online supplemental figure S2). Case-control studies (OR: 4.91; 95% CI 3.48 to 6.93, p<0.001, Z=9.03) produced a greater relative risk of AF in athletes than cohort-design studies (OR: 1.26; 95% CI 1.02 to 1.56, p=0.035, Z=2.11). However, it was not possible to appropriately analyse the association of exercise intensity and AF risk due to insufficient data.

Moderator analysis

There was a significant relationship between the mode of exercise and risk of AF (B=0.1259, p=0.0193), with mixed sport conferring a greater risk of AF compared with endurance sport (B=−0.5476, p=0.0204). Of the endurance sports, cycling conferred the highest risk of AF and Nordic skiing conferred the lowest risk (see online supplemental figure S3 for the lowest to highest risk of AF in endurance sports). Furthermore, younger (<55 years) athletes were more likely to develop AF (B=−0.02293, p<0.001). There was a significant relationship
between study design and risk of AF, with athletes in case–control studies more likely to develop AF (B = −1.3670, p < 0.001).

**DISCUSSION**

This meta-analysis aimed to quantify the risk of AF in athletes in an array of sporting activities compared with non-athlete controls. The results of the analysis indicate that the relative risk of AF is higher in athletes than in those not exercising or performing regular, non-competitive physical activity. Despite the increased risk observed among different study sizes, sport types, ages and exercise modalities, there remains a lack of high-quality studies with consistent methodologies to quantify the maximum safest regular ‘dose’ of exercise before AF risk becomes significant.

First, our OR, considering all primary studies, of 2.46 is much closer to the ‘true’ value as we included studies examining athletes, physically active individuals and sedentary populations, unlike other analyses. Additionally, the inclusion of studies involving athlete cohorts with common health conditions offers a more ‘real world’ estimation. Crucially, our research identifies a significant relationship between mode of exercise and risk of developing AF, with mixed sport conferring a greater risk than endurance sport. Of the studies included in the analysis, three involved Nordic skiing, with these studies representing 36% of the total weighting in the overall effect size. With this, it cannot be excluded that the remaining endurance sports included were comparatively underestimated in the analysis with Nordic skiing over-represented due to a greater quantity of published studies. Indeed, the observation that endurance sports correlate with AF is generally accepted. Mechanistically, Wilhelm indicates biatrial remodelling, dilation and fibrosis as crucial substrates to developing this pathology. However, Brugger et al suggested that mechanical and electrical remodelling of the atria seen with chronic endurance sport is functional and does not predispose individuals to a higher risk of AF. Despite this, current evidence does indicate a potential increase in AF risk with endurance exercise specifically. Separately, the association of ‘mixed sports’ with an increase in AF risk is difficult to interpret due to the broad range of sports analysed, complicating the process of elucidating true effects from specific training modalities. However, training volume may be an important risk factor for the development of AF and merits future research.

Our finding of a greater relative incidence of AF in younger athletes (OR: 3.60) echoes the results from Ayinde et al who performed a similar meta-analysis. Additionally, Furlanello et al examined the relationship between AF risk and sport participation in elite and Olympic athletes, finding a significant prevalence among young (average age 24 years old), male athletes. However, the OR in older athletes was still 1.76 and much higher compared with non-athletes and thus must not be ignored. Investigating the mechanistic underpinning of this age-dependant risk, Hoogsteen et al discuss AF being triggered by adrenergic surges occurring during exercise in younger athletes, whereas older athletes typically experience vagally induced AF after exercise, normally from secondary autonomic dysfunction. An interesting observation is that AF risk in the general population doubles at each decade after 50 years of age,
with a prevalence of up to 9% in 80–89 year olds.\(^8\) With this, the lower association of AF in older athletes compared with younger athletes has been previously linked to the higher rates of AF in older individuals, independent from physical activity levels.\(^2\)

Engagement with physical activity reduces CVD comorbidities that have been shown to increase the risk of AF through cumulative effects.\(^2\)

In line with our inclusion criteria (studies where athletes train at least 3 hours a week), it is clear that once over national physical activity guidelines, AF prevalence begins to rise.\(^2\)

However, it was not possible in these analyses to discern a specific duration and frequency threshold or type of exercise at which AF risk increased. Therefore, the current physical activity guidelines, whereby the benefits from the exercise outweigh the risks of any predisposition to AF, should be recommended.\(^3\)

We also aimed to ascertain AF risk in all athletes, including samples representative of those with CVD risk factors, which are independently associated with AF risk among the general population.\(^3\)

As such, we performed subanalyses on these CVD risk factors to understand their role in athlete and non-athlete AF prevalence. There was a greater prevalence of CVD in the non-athlete population, likely due to the cardiovascular protective role of physical activity; however, there was no statistically significant difference in the risk of AF in athletes and non-athletes with CVD risk factors. When CVD was excluded from analyses, the risk of AF in athletes was significantly greater compared with non-athletes (OR: 3.66).

**Study limitations**

One primary limitation of the present study is the possibility of residual confounding. The results of the included studies may be confounded in practice with studies applying different sets of adjusted variables. As detailed in Greenland and O’Rourke,\(^3\) the present analysis would have benefited from adjustment using external estimates in those studies which do not apply sufficient adjustment. Another issue with our study methodology arises around heterogeneity. Heterogeneity may develop from differences in study methodology, namely as our analyses sought to include studies of both case–control and cohort design, which account for differences in between-trial heterogeneity.\(^3\)

Our analysis was positive for publication bias when assessed using a qualitative risk of bias tool.\(^3\)

Indeed, it is possible that studies with a null hypotheses of a significant relationship between AF and athletes were not published, alongside bias in study methodologies. However, it is prudent to express diligence in assigning heterogeneity only to methodology, as differences in diagnostic criteria may change between studies, masking the ‘real’ incidence of AF.\(^3\)

Importance is placed also in observing funnel plot asymmetry for qualitative demonstration of small studies exhibiting large effects, not just for bias.\(^3\)

Indeed, 84% is well above an expected \(P\) value for a meta-analysis, probably owing to the differences in methodological parameters in the included studies.\(^3\)

However, we wanted to include all primary research undertaken with athletes to determine a more clinically relevant aspect of risk estimation for practitioners and coaching staff. This analysis may also be limited by the possibility of sparse data bias in the pooled OR estimates, as indicated by the unconventional ORs and wide confidence limits produced by some studies.

This is problematic as such bias can inflate the estimate of effect measures, which should be taken into account when interpreting the findings of the present analysis.\(^3\)

This meta-analysis contained studies using both male and female athletes and male athletes alone. Due to the limited data in female athletes, it is difficult to discern the relative risk of AF by gender. In general, risk of AF in female athletes appears to be lower than in male athletes.\(^3\)

It has been suggested that these differences in AF may be, in part, due to the proclivity for females to experience less atrial remodelling and electrocardiographic changes, alongside sex hormone differences which predispose females to higher vagal tone at rest and during exercise.\(^3\)

However, future research is required, especially given the increased female participation in elite sports in recent years. Finally, due to the lack of available data, we were unable to provide information regarding training volume and the risk of AF.

**CONCLUSION**

The risk of AF is greater in athletes compared with non-athletes. Mixed sports appear to confer a greater risk of AF. The risk of AF is relatively higher in younger athletes compared with older athletes, which is an association that should be investigated further (figure 4). Large prospective studies are needed to address current gaps in the literature including female athletes and the effect of years and volume of training.

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**Contributors** WN and JOD contributed to the conception and design of the study. WN, GP-W, JW, JE, SH, KK, MP, RS and JOD contributed to the development of the search strategy. WN, JE, SH and JOD conducted the systematic review. WN, GP-W, JE and JOD completed the acquisition of data. WN, JE, SH and JOD performed the data analysis. All authors assisted with the interpretation. WN, GP-W, JE and JOD were the principal writers of the manuscript. All authors contributed to the drafting and revision of the final article. All authors approved the final submitted version of the manuscript.

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Physical activity and risk of breast cancer, et al

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### Table S1: Adjusted status of extracted odds ratios

<table>
<thead>
<tr>
<th>Study Name</th>
<th>Adjusted status</th>
<th>Adjusted variables</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aizer et al⁷</td>
<td>Adjusted</td>
<td>Age, treatment assignment (aspirin or placebo, -carotene or placebo), BMI, diabetes, hypertension, hyperlipidaemia, parental history of MI, alcohol, smoking, fish consumption, multivitamin, vitamin C, vitamin E, LVH, CHF, CVD.</td>
</tr>
<tr>
<td>Andersen et al²</td>
<td>Adjusted</td>
<td>Age, occupation, and education level.</td>
</tr>
<tr>
<td>Baldesberger et al³</td>
<td>Unadjusted</td>
<td>N/A</td>
</tr>
<tr>
<td>Calvo et al⁴</td>
<td>Unadjusted</td>
<td>N/A</td>
</tr>
<tr>
<td>Claessen et al⁵</td>
<td>Unadjusted</td>
<td>N/A</td>
</tr>
<tr>
<td>Elosua et al⁶</td>
<td>Adjusted</td>
<td>Hypertension, age.</td>
</tr>
<tr>
<td>Karjalainen et al⁷</td>
<td>Unadjusted</td>
<td>N/A</td>
</tr>
<tr>
<td>Molina et al⁸</td>
<td>Adjusted</td>
<td>Age, systolic and diastolic blood pressure.</td>
</tr>
<tr>
<td>Mont et al⁹</td>
<td>Unadjusted</td>
<td>N/A</td>
</tr>
<tr>
<td>Myrstad et al¹⁰</td>
<td>Adjusted</td>
<td>CHD, hypertension, diabetes, age, height, BMI, education level, alcohol, smoking habits, leisure time physical activity.</td>
</tr>
<tr>
<td>Myrstad et al¹¹</td>
<td>Adjusted</td>
<td>Age, sex, BMI, concomitant CVD, stroke, diabetes, number of race participations.</td>
</tr>
<tr>
<td>Schreiner et al¹²</td>
<td>Unadjusted</td>
<td>N/A</td>
</tr>
<tr>
<td>Van Buuren et al¹³</td>
<td>Unadjusted</td>
<td>N/A</td>
</tr>
</tbody>
</table>

Note: BMI, body mass index; MI, myocardial infarction; LVH, left ventricular hypertrophy; CHF, congestive heart failure; CVD, cardiovascular disease; CHD, coronary heart disease; N/A, not applicable.
Where ORs were not reported, they were calculated via the CMA in-built algorithm below:

- A = Developed AF Athletes
- B = Developed AF Controls
- C = Did Not Develop AF Athletes
- D = Did Not Develop AF Controls

\[
\text{LogOddsRatio} = \log\left(\frac{A \times D}{B \times C}\right) \\
\text{LogOddsVariance} = \left(\frac{1}{A} + \frac{1}{B} + \frac{1}{C} + \frac{1}{D}\right) \\
\text{LogOddsSE} = \sqrt{\text{LogOddsVariance}} \\
\text{OddsRatio} = \exp(\text{LogOddsRatio})
\]

\[
\text{LogOddsRatio} = \log\left(\frac{83 \times 72}{24 \times 37}\right) = 1.907 \\
\text{LogOddsVariance} = \left(\frac{1}{83} + \frac{1}{24} + \frac{1}{37} + \frac{1}{72}\right) = 0.095 \\
\text{LogOddsSE} = \sqrt{0.095} = 0.308 \\
\text{OddsRatio} = \exp(1.907) = 6.730
\]
Table S2: Meta-analysis output table displaying weightings, odds ratios, fixed and random effects.

<table>
<thead>
<tr>
<th>Study Name</th>
<th>Odds Ratio</th>
<th>Lower Limit</th>
<th>Upper Limit</th>
<th>Z Value</th>
<th>P value</th>
<th>Weight (random)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aizer et al¹</td>
<td>1.200</td>
<td>1.021</td>
<td>1.411</td>
<td>2.207</td>
<td>0.027</td>
<td>12.976</td>
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<td>Andersen et al²</td>
<td>1.200</td>
<td>0.930</td>
<td>1.549</td>
<td>1.399</td>
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<td>3.880</td>
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<td>Elosua et al⁶</td>
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<td>Mont et al⁹</td>
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Figure S1: Study quality funnel plot with standard error vs logarithmic odds ratio.

<table>
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<th>Quality</th>
<th>Odds Ratio (95% CI)</th>
<th>Z-Value</th>
<th>p-Value</th>
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<td>Good Quality</td>
<td>2.322 (1.623 – 3.322)</td>
<td>4.610</td>
<td>&lt;0.001</td>
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<tr>
<td>Poor Quality</td>
<td>4.798 (1.219 – 18.878)</td>
<td>2.244</td>
<td>0.025</td>
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<tr>
<td>Overall</td>
<td>2.432 (1.720 – 3.439)</td>
<td>5.028</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

Figure S2: Subgroup analysis of study quality and the risk of atrial fibrillation.
Figure S3: Endurance sport meta-regression bubble plot outputs.

Regression of Log odds ratio on Endurance Sports
References