Atrial functional and geometrical remodeling in male highly trained athletes - for better or worse?

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Word count: 3395
Abstract

Purpose: Highly trained athletes have an increased risk of atrial arrhythmias. Atrial geometrical and functional remodeling may be the underlying substrate. We analyze and relate atrial size, deformation and performance in professional handball players compared with non-sportive subjects.

Methods: 24 professional handball players and 20 non-sportive males were compared. All subjects underwent an echocardiographic study with evaluation of left (LA), right atrial (RA) dimensions and deformation by strain (Sa) and strain rate (SRa). Atrial performance was assessed from the atrial stroke volume (SV). With computational geometrical models, we studied the relation between atrial volumes, strains and SV and compared atrial working conditions. We estimated the functional reserve and a resulting average wall stress.

Results: LA and RA volumes were larger in athletes than in controls (35.2±8.8 vs 24.8±4.3ml/m², p<0.01 and 29.0±8.4 vs 19.0±5.1ml/m², p<0.01 respectively). LASa and RASa during active atrial contraction were decreased in athletes (-12.2±2.0 vs -14.5±2.1%, p<0.01 and -12.1±1.8 vs -14.2±1.5%, p<0.01 respectively). LASV was similar between groups (6.6±1.4 vs 7.3±1.1 ml, p=0.19) and RASV was lower in athletes (6.2±1.3 vs 7.2±1.1 ml, p<0.01).

Computational models showed that this different operational mode potentially increases performance reserve, but at the cost of higher atrial wall stress.

Conclusion: a proportion of athletes with enlarged LA and RA showed different atrial contractile performance, likely resulting in atria working at higher wall stress.

Key words: atrial strain; atrial function; echocardiography; wall stress; exercise.
Abbreviations

AF, atrial fibrillation
CMR, cardiac magnetic resonance
LA, left atrium
LASa, peak negative left atrial strain during active atrial contraction
LASRa, peak negative left atrial strain rate during active atrial contraction
LASRe, LA early diastolic peak strain rate
LASRs, LA peak strain rate during ventricular systole
LASs, LA peak strain during ventricular systole
LV, left ventricle
RA, right atrium
RASa, peak negative right atrial strain during active atrial contraction
RASRa, peak negative right atrial strain rate during active atrial contraction
RASRe, RA early diastolic peak strain rate
RASRs, RA peak strain rate during ventricular systole
RASs, RA peak strain during ventricular systole
RV, right ventricle
SV, stroke volume
Introduction

Observational studies have reported an increased prevalence of atrial fibrillation (AF) in highly trained athletes, as compared to case-matched controls from the general population [Elosua et al. 2009 and Molina et al. 2008]. Particularly, a cohort of retired handball players showed an AF incidence of 30% at the age of 50 [Van Buuren F et al. 2012]. A meta-analysis of these observational studies showed a five-fold increase in the overall risk of AF in this population [Abdulla J and Rokkedal J. 2009]. The etiology of AF in this particular group of subjects is still poorly understood and several theories have been suggested [Turagam M et al. 2012]. Increased vagal tone with consequent bradycardia may lead to dispersion of atrial repolarization, which in turn may increase the susceptibility to AF [Wilhelm M et al. 2011]. The remodeling process induced in the atria by intensive exercise, together with the above factors, may translate into a potential dysfunction [Pellicia A et al. 2005], and lead to the development of the arrhythmia.

Atrial flutter can also be present or co-exist with AF and has been linked to right atrial dilatation in the setting of the athlete’s heart [Erol MK and Karakelleoglu S. 2002].

In recent years, atrial strain and strain-rate analysis by two-dimensional speckle tracking has emerged as a novel method to evaluate left atrial (LA) and right atrial (RA) function [Padelleti M et al. 2012; Saraiva RM et al. 2010]. The assessment of LA geometry [Tsang MY et al. 2012], and function by strain and strain-rate has been used as a predictor of the occurrence of AF in different clinical scenarios such as pulmonary veins ablation [Schneider C et al. 2008], and coronary artery by-pass grafting [Gabrielli L et al. 2009]. Additionally, it was shown that increased local wall stress is associated with focal atrial remodeling and electrical alterations [Hunter RJ et al. 2012].
Simple computational models can help understanding the link between chamber size, deformation and contractile performance [Bijnens BH et al. 2012]. In the present study, we apply an ellipsoid-based computational model to the data collected from a set of athletes and control subjects, in order to reach a better understanding of how these variables interrelate in these populations. This model has been previously used to assess ventricular wall stress and to perform a detailed analysis of ventricular performance [Mirsky et al. 1981]. The objective of the present study is to provide insights into the atrial remodeling process in athletes; our hypothesis is that a subgroup of athletes might have different atrial working conditions that might favour the development of a potential arrhythmogenic substrate.

Methods

Study design and population

This was a cross-sectional study, comparing two groups: 24 athletes and 20 sedentary healthy controls with similar age who were recruited among hospital staff and patients’ relatives with a strictly normal medical record. All participants were male, in sinus rhythm, normotensive, did not take any medication or anabolic drug and did not have other cardiovascular disease or risk factors. The athletes group consisted of professional handball players from the same national team. All of them had been training an average of 15 hours/week during the past year. All subjects in the control group did not participate in routine competitive or recreational sports.

The study was approved by the ethics committee of our institution and informed consent was obtained from all individuals.
Echocardiography

Each individual underwent a complete two-dimensional echocardiographic study using a commercially available ultrasound scanner (Vivid Q, GE Healthcare, Milwaukee, WI) with a 2.5-MHz phased array transducer (M4S). Standard echocardiographic views were obtained. Images were analyzed off-line with a commercially available software (EchoPac version 108.1.6, GE Healthcare). Left ventricle (LV) and right ventricle (RV) measurements were done according to the recommendations of the American Society of Echocardiography [Lang RM et al. 2006]. LA volume was calculated from apical four and two-chamber views using the biplane method of discs and RA volume was calculated from a monoplane derived volume (apical four chamber view). LA and RA emptying fractions were calculated using the difference between the maximal volumes (just before the opening of the mitral valve) and the minimal ones (at the closure of the mitral valve) / maximal volume. Atrial active stroke volume (SV) was estimated using the difference between the pre-A volume (obtained from the last frame before mitral valve opening or at time of P wave on the surface electrocardiogram) and minimal volume. Also, from the measured atrial volumes we calculated LA/RA volumes of early diastolic phase (maximal volume – pre-A volume) and LA/RA emptying fractions of early diastolic phase ([maximal volume – pre-A volume]/maximal volume). Area and volume measurements were indexed by body surface area to allow proper comparison between groups and all data were analyzed by two blinded, experienced sonographers.

To define severe atrial dilatation, a cut-off of 40 ml/m² was used for both atria [Lang RM et al. 2006].

Atrial strain and strain rate
Strain and strain rate of both atria were analyzed off-line with a commercially available software package (2Dstrain, EchoPac version 108.1.6, GE Healthcare). Acquisitions were performed with an optimized image with a frame rate of at least 50 fps and with focal point in atriums. The endocardial border was manually traced using a point-and-click technique. LA and RA strains were calculated with the reference point set at the onset of the P-wave of the surface ECG, which allowed identifying the peak negative strain (shortening) during atrial contraction (LASa and RASa, respectively) Similarly, the peak negative strain rate (active shortening) during atrial contraction was identified (LASRa and RASRa, respectively). The software divided the atrial wall into 6 segments and the average was taken for analysis (Figures 1A and 1B). Also, LA/RA peak strain and strain rate during ventricular systole and strain rate during early diastolic phase were recorded.

**Computational geometrical model**

The possible relations between atrial volume (pre-A volume, taken before the onset of atrial contraction), the global atrial longitudinal strain during active contraction, and the SV generated by the atrial contraction, were investigated with a simple geometrical mathematical model.

In this model, the atria were approximated by ellipsoids with two of the (short) radii being equal and with long axis / short axis ratios of 1.26 (left atrium) and 1.16 (right atrium), corresponding to the average values measured from our data. In order to simulate the potential range within which the atria of the studied subjects could function, we calculated the volume-strain-SV relations for a pre-A volume from 18 to 95 ml (for the LA) or 19 to 105 ml (for the RA), and for a SV from 3.5 to 10 ml (for both atria). The global strain during atrial contraction was calculated from the pre-A and minimal volumes based on the radius change during atrial contraction. Global
atrial longitudinal strain was computed as the change in the largest circumference, whose formula is detailed in Appendix.

Next, in order to find the (resting) working point in this volume-strain-SV relation for each individual from our study, the ellipsoid model was fitted on each of the individual’s measured atrial volumes and diameters, as also detailed in Appendix. This personalized model provided an estimation of the SV reserve and a rough approximation of average atrial wall stress for each individual.

**Statistical methods**

Continuous baseline variables were expressed as mean ± standard deviation (SD) values or median (interquartile range), after checking for normal distribution as assessed by the Kolmogorov-Smirnov test. Categorical variables were expressed as total number (percentages) and compared between groups using Chi-square or Fisher’s test when appropriate. Continuous variables were tested by unpaired t test or Mann-Whitney U test, according to normality, and paired data by paired t test or Wilcoxon analysis. Pearson or Spearman methods were used to analyze the correlation between continuous variables when suitable. Reproducibility of the observations was assessed offline using Bland-Altman analysis on 10 randomly selected subjects, from 2 different times (intra-observer) and by 2 different observers (inter-observer). The acquisitions were performed by one, blind expert operator and all the measurements were performed from the same acquisition. Statistical significance was established at P<0.05. All data were analyzed using the SPSS statistical package version 15.0 (SPSS Inc., Chicago, IL).
Results

Population characteristics
The clinical characteristics of the studied population are listed in Table 1. All participants were male. Body mass index and body surface area were significantly higher in athletes than in controls. In all subjects, no significant mitral or tricuspid regurgitation were found.

Table 2 shows the characteristics of LA and LV in athletes and controls. LV dimensions and function were similar in both groups while LV mass was higher in athletes. LA maximum volume was increased in athletes as compared to controls, LA emptying fraction and active LA SV (both volumetric and Doppler-based) were similar between both groups and athletes showed reduced LASa and LASRa as compared to controls.

Table 3 shows RV and RA dimensions and function. Athletes had increased RV basal diameter as compared to controls and RV function was similar between groups. In athletes, RA maximum volume was increased and RASa and RASRa were reduced as compared to controls.

We did not find differences in the other phases of LA/RA cycle regarding emptying fraction and deformation (Table 2 and 3).

Atrial size, deformation and contractile performance
Figures 2A, 2B and 2C show the relation between the active LA deformation and the maximal LA atrial size, for all individuals. A proportion of athletes (circle in the figure) showed a larger LA volume, lower deformation (LASa and LASRa) and lower LA SV during active atrial contraction: athletes with LA volume ≥ 40 ml/m² (n=7) showed significantly reduced LASa, LASRa and active LA SV during atrial contraction as compared to athletes with LA volume < 40 ml/m².
Similarly to what was reported for the LA, Figures 2D, 2E and 2F show the relation between RA maximal volume and RASa and RA SV. A proportion of athletes (circle in the figure) also showed a larger RA volume, lower deformation (RASa and RASRa) and lower RA SV during active atrial contraction, and similarly to what was observed with LA: athletes with a RA volume ≥ 40 mL/m² (n = 4) showed significantly reduced RASa, RASRa and RA SV during active atrial contraction as compared to athletes with RA volume < 40 mL/m² (-9.55 ± 0.92% vs -12.63 ± 1.76%, p < 0.01 [RASa]; -0.99 ± 0.07 s⁻¹ vs -1.38 ± 0.53 s⁻¹, p = 0.04 [RASRa] and 4.7 ± 0.7 mL vs 6.5 ± 1.2 mL, p =0.01 [RA SV]). All athletes with severe RA dilatation showed severe LA dilatation.

Among athletes there was a significant correlation between LASa with LA maximal volume (r = 0.57, p < 0.01), with LASRa (r = 0.73, p < 0.01) and with LA SV (r = - 0.63, p < 0.01); and between RASa with RA maximal volume (r = 0.64, p < 0.01) with RASRa (r=0.62, p < 0.01) and with RA SV (r = - 0.78, p < 0.01).

Finally, athletes showed a significant correlation between LASa and RASa and between LA and RA volume (r = 0.50, p = 0.017 [strain]; r = 0.73, p < 0.01 [volumes]), indicating a similar performance/remodeling of both atria in a given athlete.

**Atrial strain and strain rate reproducibility**

The inter-observer and intra-observer agreement for LA and RA strain and strain rate are detailed in Table 4. In summary, the inter-observer variability was < 6% and the intra-observer variability was < 2%.
Geometrical modeling

As commented in the Methods section, an ellipsoid-based computational model was used for (i) interpreting of the relation between the above-measured parameters (atrial size, deformation and contractile performance), and (ii) for investigating the link between atrial SV reserve, wall stress, and volume.

Figure 3 shows the expected (geometrical) relations between deformation, size and SV (solid lines) for different combinations of pre-A volume and SV in the LA and RA. The working point of each of individual (athletes and controls) was specified (colored dots). The figure indicates that individuals with larger volumes need a lower strain to maintain a similar SV. Additionally, while several athletes were located among the control subjects, a subset of athletes had larger atria with lower strain and SV. Observations were comparable for both left and right atria.

Figure 4 represents the predicted atrial SV reserve (top), the maximal atrial wall stress (middle) and the atrial wall stress associated with a predicted atrial SV reserve (bottom), for each of the individuals in the study. Note that these data results from the personalization of the model to each individual’s data, as commented in the Methods section. As it can be observed, a majority of the athletes had a larger SV reserve because of working with larger atria (and therefore lower strain at rest, as shown in figure 3). However, this comes at the cost of elevated wall stress. Additionally, different subgroups of athletes may be considered. The ones with large atrial volumes present a high potential to increase the SV, but at the cost of a high wall stress. The second subgroup has a size similar to controls, with a moderate potential to increase SV, but with apparently higher wall stress than controls because of a different aspect ratio of the atrium. Observations were comparable for both left and right atria.
Discussion

Our study first showed that, in a population of young athletes (professional elite handball players), both atria have larger volumes and reduced deformation during active atrial contraction at rest as compared to controls, and a subgroup of athletes has significantly higher volumes and lower deformation. Then, the use of an ellipsoid-based computational model allowed understanding the different working conditions of these subgroups. Larger volumes provide an acute improved performance (by requiring lower deformation to maintain a similar stroke volume) and reserve, but at the cost of higher wall stress. Our interpretation is that this might lead to long term damage, which could explain the increased incidence of atrial arrhythmia observed in this population.

Structural remodeling is seen as the main contributor for initiation and persistence of AF or atrial flutter [Sitges M et al. 2009]. It has been shown that the LA may be enlarged in as many as 20% of competitive athletes [Pelliccia A et al. 2005], but not all of these subjects developed AF. Indeed, other mechanisms, besides increased atrial size, have been proposed to explain the development of an arrhythmia, such as increased parasympathetic tone, reduced sympathetic tone and increased inflammation [Sorokin AV et al. 2011].

While LA enlargement is common in highly trained athletes, it is suggested that this represents a physiologic adaptation to exercise and chronic training and not a consequence of increased left ventricular filling pressures secondary to diastolic dysfunction [D’Andrea A et al. 2010]; accordingly, the risk of atrial arrhythmias is mildly elevated and developed later in life [Molina L et al 2008; Van Buuren F et al 2012]. Our results confirm that resting LV filling pressures on the
basis of E/e’ ratio and diastolic function are not different in these highly trained athletes and the control group.

In an animal model of chronic endurance training, it was demonstrated that additionally to atrial enlargement, collagen content and myocardial fibrosis markers were significantly increased in both atria after training; these histological and molecular findings were observed together with an increased inducibility of atrial arrhythmias [Benito B et al. 2011]. In the present study we could see that, while a majority of athletes showed atrial properties similar to controls, there was a subgroup presenting marked bi-atrial remodeling. These athlete’s atria are working at lower strain and larger volumes for a given or lower, atrial stroke volume as compared to controls. While this might be beneficial to have a larger SV reserve (since when increasing the strain, with a larger atrium they could generate much more SV), it would also result in higher wall stress. Since there is an intrinsic relation between size, deformation and SV, and that individuals can perform at different ‘working points’, this does not necessarily imply that larger dilatation is a synonym for atrial dysfunction (it is unclear whether this dilatation is indeed physiological). It could be the result of having to cope with the higher SV demand in better performing athletes, or it could also be dilatation and the decrease in deformation and contractility as a result of ongoing atrial wall damage, similar to what can be seen in severe valve regurgitation [Margreet A et al. 2010], or other overload situation [Marciniak A et al. 2010]. Nonetheless, it is clear that the elevated wall stress is an important risk factor for the initiation/worsening of myocardial damage, and that on the longer run, this will result in increased vulnerability to atrial arrhythmias [Di Martino E et al. 2011; Tsai CT et al. 2011].

Consequently, in order to determine whether the atria in an individual are dysfunctional, a more profound analysis of deformation, size and performance, and especially the response to exercise
(testing if the predicted SV increase can indeed be realized), together with some kind of tissue
coloration (e.g. using CMR delayed enhancement), is required [Prinz C et al. 2012]. This
means that the understanding of the direct link between atrial remodeling and the risk of
arrhythmia in athletes still needs to be improved.

In our study we also discovered a significant correlation between LASa and RASa showing that
both atria of a given individual receive similar remodeling triggers, which also could explain the
concomitant occurrence of LA and RA arrhythmias in those athletes with pathologic atrial
remodeling [Hoogsteen J et al. 2004].

Regarding the other phases of the atrial cycle we did not find differences in terms of emptying
fraction and deformation in agreement with other previous report [Gabrielli et al. 2012].

One limitation of our work is that our model only provides an average approximation of atrial
wall stress based on global morphologic characteristics. Given the complex shape of the atria, a
3D localized approach would be needed to get accurate estimates of wall stress. However, the
estimated values are in range of what has been published before for a detailed analysis, and since
we are only interested in the tendency for wall stress rather than the exact values or the individual
comparison, they provide an acceptable first approximation. The model-based calculation of SV
reserve assumes a normal, physiologic response to exercise and does not include abnormal or
pathological changes in atrial function (e.g. inability to increase deformation and thus SV). While
the precision of such a modeling might be discussed, this approximation allows us to identify
subgroups with different behaviors. Other important limitations are the small sample size and the
fact that these findings were not extended to other sports disciplines or to a female population.

Current speckle tracking tools may perform worse when applied to the right side of the heart and
the atria, due to the potential lower quality of the images (specially considering the worse lateral
resolution in 2-dimensional echocardiography) and the fact that the algorithms are developed for
the left ventricle. However, we believe these measurements are reliable, because an adequate
endocardial tracking is ensured and checked frame-by-frame and deformation values are in
keeping with what is seen in the image.

Finally, athlete’s atrial volumes are not particularly high (D’Andrea et al. 2013) but we used the
proposed cut off value of 40 ml/m proposed by guidelines as severe dilatation (Lang et al. 2006)
to separately analyze extreme values of atrial size.

Even considering all the limitations, we believe that our findings, especially given the highly
comparable group of individuals, are robust enough to generate a hypothesis to be confirmed in
larger studies and including a more elaborated analysis of atrial function, in particular during
exercise.

Conclusion

To our knowledge this is the first study that assessed active atrial function in highly trained
athletes and showed that some of these show remodeling that is suggestive of atrial dysfunction.

Why some athletes from the same team, with similar training and thus comparable cardiac
overload, show completely different adaptation/remodeling, remains an unanswered question and
might be related to other predisposing factors that increase the risk of pathologic atrial
remodeling and increase the vulnerability to atrial arrhythmias.
Acknowledgements

This study was partially supported by grants from the "Generalitat de Catalunya (Consell Catala
de l’Esport)”, from "Grupo Memora (SFB-Grupo Memora)”, Spanish Society of Cardiology
(Spanish Heart Foundation), from the “Plan Nacional I+D+I”, Spanish Government (DEP2010-
20565), and from the "Subprograma de Proyectos en Salud”, Instituto de Salud Carlos III,
Spanish Government (FIS - PI11/01709).

Authors have nothing to disclose
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Cardiovascular Research 89:805-815.


Table 1. Clinical characteristics of the studied subjects.

<table>
<thead>
<tr>
<th></th>
<th>Controls (n = 20)</th>
<th>Athletes (n = 24)</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>27 ± 4</td>
<td>28 ± 4</td>
<td>NS</td>
</tr>
<tr>
<td>Weight (kg)</td>
<td>72 ± 8</td>
<td>94 ± 12</td>
<td>&lt; 0.01</td>
</tr>
<tr>
<td>Height (cm)</td>
<td>175 ± 15</td>
<td>191 ± 15</td>
<td>&lt; 0.01</td>
</tr>
<tr>
<td>BMI (kg/m²)</td>
<td>23 ± 2.7</td>
<td>25 ± 2.6</td>
<td>0.01</td>
</tr>
<tr>
<td>BSA (m²)</td>
<td>1.80 ± 0.12</td>
<td>2.20 ± 0.17</td>
<td>&lt; 0.01</td>
</tr>
<tr>
<td>Heart rate (bpm)</td>
<td>76 ± 12</td>
<td>55 ± 9</td>
<td>&lt; 0.01</td>
</tr>
</tbody>
</table>

Abbreviations: BMI, body mass index; BSA, body surface area

NS, non-significant statistical difference.
Table 2. Echocardiographic left ventricular and left atrial characteristics of the studied subjects.

<table>
<thead>
<tr>
<th></th>
<th>Controls (n = 20)</th>
<th>Athletes (n = 24)</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>LV End-diastolic diameter (cm)</td>
<td>5.1 ± 0.3</td>
<td>5.3 ± 0.6</td>
<td>NS</td>
</tr>
<tr>
<td>LV mass index (g/m²)</td>
<td>86 ± 10</td>
<td>116 ± 21</td>
<td>&lt; 0.01</td>
</tr>
<tr>
<td>LV ejection fraction (%)</td>
<td>60 ± 6</td>
<td>57 ± 5</td>
<td>NS</td>
</tr>
<tr>
<td>LV GLS (%)</td>
<td>-20.0 ± 1.3</td>
<td>-19.0 ± 1.9</td>
<td>NS</td>
</tr>
<tr>
<td>LA anteroposterior diameter (cm)</td>
<td>3.5 ± 0.2</td>
<td>4.1 ± 0.3</td>
<td>&lt; 0.01</td>
</tr>
<tr>
<td>LA area index (cm²/m²)</td>
<td>8.6 ± 1.2</td>
<td>10.1 ± 1.3</td>
<td>&lt; 0.01</td>
</tr>
<tr>
<td>LA volume index (ml/m²)</td>
<td>24.8 ± 4.3</td>
<td>35.2 ± 8.8</td>
<td>&lt; 0.01</td>
</tr>
<tr>
<td>LA pre A volume index (mL/m²)</td>
<td>12.9 ± 3.4</td>
<td>19.6 ± 9.2</td>
<td>&lt; 0.01</td>
</tr>
<tr>
<td>LA minimal volume index (ml/m²)</td>
<td>9.5 ± 3.4</td>
<td>16.6 ± 8.5</td>
<td>&lt; 0.01</td>
</tr>
<tr>
<td>LA emptying fraction</td>
<td>0.59 ± 0.23</td>
<td>0.61 ± 0.35</td>
<td>NS</td>
</tr>
<tr>
<td>LA emptying fraction in early diástole</td>
<td>0.45 ± 0.09</td>
<td>0.46 ± 0.15</td>
<td>NS</td>
</tr>
<tr>
<td></td>
<td>Group 1</td>
<td>Group 2</td>
<td>p-Value</td>
</tr>
<tr>
<td>------------------------------</td>
<td>---------------</td>
<td>---------------</td>
<td>----------</td>
</tr>
<tr>
<td>LA A wave VTI (cm)</td>
<td>4.9 ± 1.0</td>
<td>4.6 ± 0.7</td>
<td>NS</td>
</tr>
<tr>
<td>LA active contraction SV (ml)</td>
<td>7.3 ± 1.1</td>
<td>6.6 ± 1.4</td>
<td>NS</td>
</tr>
<tr>
<td>LA emptying volume in early</td>
<td>21.5 ± 6.2</td>
<td>34.8 ± 10.2</td>
<td>&lt; 0.01</td>
</tr>
<tr>
<td>diastole (mL)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>E wave velocity (cm/s)</td>
<td>78 ± 13</td>
<td>79 ± 14</td>
<td>NS</td>
</tr>
<tr>
<td>A wave velocity (cm/s)</td>
<td>37 ± 13</td>
<td>42 ± 7</td>
<td>NS</td>
</tr>
<tr>
<td>EA ratio</td>
<td>2.1 ± 0.6</td>
<td>1.9 ± 0.4</td>
<td>NS</td>
</tr>
<tr>
<td>E wave deceleration time (ms)</td>
<td>178 ± 47</td>
<td>205 ± 51</td>
<td>NS</td>
</tr>
<tr>
<td>A wave duration (ms)</td>
<td>133 ± 15</td>
<td>143 ± 23</td>
<td>NS</td>
</tr>
<tr>
<td>Em septal (cm/s)</td>
<td>9.8 ± 1.6</td>
<td>9.2 ± 1.8</td>
<td>NS</td>
</tr>
<tr>
<td>Em lateral (cm/s)</td>
<td>13.2 ± 2.6</td>
<td>13.1 ± 2.1</td>
<td>NS</td>
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<tr>
<td>E/Em ratio</td>
<td>7.4 ± 1.1</td>
<td>7.0 ± 1.0</td>
<td>NS</td>
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<tr>
<td>LA Sa (%)</td>
<td>-14.5 ± 2.1</td>
<td>-12.2 ± 2.0</td>
<td>&lt; 0.01</td>
</tr>
<tr>
<td>LASRa (s⁻¹)</td>
<td>-1.87 ± 0.36</td>
<td>-1.52 ± 0.59</td>
<td>0.02</td>
</tr>
<tr>
<td>LASs (%)</td>
<td>36.2 ± 6.0</td>
<td>35.9 ± 5.6</td>
<td>NS</td>
</tr>
<tr>
<td>LASRe (s⁻¹)</td>
<td>-2.15 ± 0.59</td>
<td>-2.18 ± 0.47</td>
<td>NS</td>
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<tr>
<td>LASRs (s⁻¹)</td>
<td>1.65 ± 0.48</td>
<td>1.64 ± 0.40</td>
<td>NS</td>
</tr>
</tbody>
</table>
Abbreviations: Em, early peak velocity of the mitral annulus by Doppler Tissue Imaging; GLS, global longitudinal strain; LA, left atrium; LASa, LA late diastolic (atrial contraction) peak strain; LASRa, LA late diastolic (atrial contraction) peak strain rate; LASRe, LA early diastolic peak strain rate; LASRs, LA peak strain rate during ventricular systole; LASs, LA peak strain during ventricular systole; LV, left ventricle; NS, non-significant statistical difference; SV, stroke volume (planimetry).
Table 3. Echocardiographic right ventricular and right atrial characteristics of the studied subjects.

<table>
<thead>
<tr>
<th></th>
<th>Controls (n = 20)</th>
<th>Athletes (n = 24)</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>RV basal diameter (mm/m²)</td>
<td>16.0 ± 3.1</td>
<td>17.0 ± 4.1</td>
<td>0.05</td>
</tr>
<tr>
<td>RV fractional area change (%)</td>
<td>40 ± 5</td>
<td>38 ± 4</td>
<td>NS</td>
</tr>
<tr>
<td>TAPSE (mm)</td>
<td>28 ± 3</td>
<td>27 ± 2</td>
<td>NS</td>
</tr>
<tr>
<td>RV GLS (%)</td>
<td>-23.7 ± 3.3</td>
<td>-22.8 ± 3.3</td>
<td>NS</td>
</tr>
<tr>
<td>RA maximal volume index (ml/m²)</td>
<td>19.0 ± 5.1</td>
<td>29.0 ± 8.4</td>
<td>&lt; 0.01</td>
</tr>
<tr>
<td>RA pre A volume index (ml/m²)</td>
<td>14.3 ± 4.1</td>
<td>20.9 ± 8.6</td>
<td>&lt; 0.01</td>
</tr>
<tr>
<td>RA minimal volume index (ml/m²)</td>
<td>9.8 ± 3.0</td>
<td>18.1 ± 7.9</td>
<td>&lt; 0.01</td>
</tr>
<tr>
<td>RA active contraction SV (ml)</td>
<td>7.2 ± 1.1</td>
<td>6.2 ± 1.3</td>
<td>&lt; 0.01</td>
</tr>
<tr>
<td>RA emptying volume in early diastole (mL)</td>
<td>11.0 ± 7.6</td>
<td>18.0 ± 10.8</td>
<td>0.02</td>
</tr>
<tr>
<td>RA emptying fraction</td>
<td>0.48 ± 0.15</td>
<td>0.39 ± 0.16</td>
<td>NS</td>
</tr>
<tr>
<td>RA emptying fraction in early diastole</td>
<td>0.31 ± 0.17</td>
<td>0.30 ± 0.17</td>
<td>NS</td>
</tr>
<tr>
<td>RASa (%)</td>
<td>-14.2 ± 1.5</td>
<td>-12.1 ± 1.8</td>
<td>&lt; 0.01</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Value 1</td>
<td>Value 2</td>
<td>p-Value</td>
</tr>
<tr>
<td>-------------</td>
<td>---------</td>
<td>---------</td>
<td>---------</td>
</tr>
<tr>
<td>RASRa (s⁻¹)</td>
<td>-2.05 ± 0.48</td>
<td>-1.32 ± 0.37</td>
<td>&lt; 0.01</td>
</tr>
<tr>
<td>RASs (%)</td>
<td>34.3 ± 7.5</td>
<td>33 ± 7.1</td>
<td>NS</td>
</tr>
<tr>
<td>RASRe (s⁻¹)</td>
<td>-1.98 ± 0.52</td>
<td>-1.68 ± 0.50</td>
<td>NS</td>
</tr>
<tr>
<td>RASRs (s⁻¹)</td>
<td>2.00 ± 0.51</td>
<td>1.82 ± 0.35</td>
<td>NS</td>
</tr>
</tbody>
</table>

Abbreviations: GLS, global longitudinal strain; NS, non-significant statistical difference; RA, right atrium; RASa, RA late diastolic (atrial contraction) peak strain; RASRa, RA late diastolic (atrial contraction) peak strain rate; RASRe, RA early diastolic peak strain rate; RASRs, RA peak strain rate during ventricular systole; RASs, RA peak strain during ventricular systole; RV, right ventricle; SV, stroke volume (planimetry); TAPSE, tricuspid annulus plane systolic excursion.
Table 4. Inter- and intraobserver reproducibility of the speckle tracking measurements

<table>
<thead>
<tr>
<th></th>
<th>Interobserver agreement</th>
<th>Intraobserver agreement</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean difference</td>
<td>CI 95%</td>
</tr>
<tr>
<td>LASe (%)</td>
<td>0.4</td>
<td>-1.90 to 2.60</td>
</tr>
<tr>
<td>LASe (s⁻¹)</td>
<td>-0.05</td>
<td>-0.50 to 0.60</td>
</tr>
<tr>
<td>RASe (%)</td>
<td>-0.8</td>
<td>-3.20 to 1.60</td>
</tr>
<tr>
<td>RASe (s⁻¹)</td>
<td>-0.10</td>
<td>-0.57 to 0.37</td>
</tr>
</tbody>
</table>

Abbreviations: CI, confidence interval; LASe, LA late diastolic (atrial contraction) peak strain; LASe, LA late diastolic (atrial contraction) peak strain rate; RASe, RA late diastolic (atrial contraction) peak strain; RASe, RA late diastolic (atrial contraction) peak strain rate.
Figures title and legend

**Figure 1.** Left atrial strain (A) and strain rate (B) curves obtained from 2D speckle tracking. Arrows indicate LASa and LASRa, respectively.

**Figure 2.** Relation between the active LA/RA deformation and maximal atrial size in athletes and controls.

Top panels circle: subgroup of athletes with higher LA dilatation and lower LASa (2A), LASRa (2B) and LA active SV by planimetry (2C).

Bottom panels circle: subgroup of athletes with higher RA dilatation and lower RASa (2D), RASRa (3E) and RA active SV by planimetry (3F).

**Figure 3.** Modeled relation between pre-A volume, strain and SV with individual data overlaid.

**Figure 4.** SV reserve to achieve 40% strain (top); stress-volume relation (middle and bottom). The curve shows the theoretical stress-volume relation, calculated according to equation [27].
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Figure 1
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Figure 2
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A) LA volume index (mL/m²)

B) LA SV (mL)

C) LA SVa (L/min)

D) RA volume index (mL/m²)

E) RA SV (mL)

F) RA SVa (L/min)
Figure 3
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NewFigure4
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