#### **ORIGINAL PAPER**



# Influence of different dynamic sporting disciplines on right ventricular Structure and function in elite male athletes

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#### Abstract

Our objective was to assess the influence of different levels of exposure to dynamic training on right ventricular (RV) structure, function and mechanics in elite male athletes. We recruited 492 male elite athletes aged between 18 and 30 years old. Athletes were grouped according to their sporting discipline using the Mitchell Classification as Low Dynamic (LD), Moderate Dynamic (MD) or High Dynamic (HD). All participants underwent 2D, Doppler, tissue Doppler and strain ( $\epsilon$ ) echocardiography with a focused and comprehensive assessment of the right heart. Athletes involved in MD sports had the largest absolute RV chamber size and when scaled to body size RVOT PLAX, RVOT2, RVD1 and RVD3 were larger in HD compared to MD and LD athletes. There were no between group differences in conventional RV functional indices as well as global RV  $\epsilon$  (LD:  $-23.4\pm3.1$  vs. MD:  $-22.7\pm2.7$  vs. HD:  $-23.5\pm2.6$ , %) and strain rate (P>0.01). The base to apex  $\epsilon$  gradient in the RV septum was lower in the MD athletes compared to HD and LD due to a lower apical septal  $\epsilon$  which significantly correlated with absolute RV chamber size. After scaling for body size there was evidence of larger RV cavities in both MD and HD athletes compared to LD athletes. Global RV function,  $\epsilon$  and strain rate were not different between groups. MD athletes had lower apical septal  $\epsilon$  that contributed to a smaller base-to-apex  $\epsilon$  gradient that is partially associated with larger absolute RV chamber dimensions.

Keywords Right ventricule · Athlete's heart · Strain · Echocardiography · Exercise

# Introduction

The term athlete's heart (AH) is used to describe the physiological cardiac adaptation in response to exercise training [1–4]. Many studies of AH have focused on the left ventricle but the right ventricle (RV) also undergoes physiological remodelling due to the hemodynamic overload associated with training [2, 5–7]. The nature of RV remodelling and functional change associated with prolonged training exposure often leads to a phenotypical presentation beyond normal limits that can create a diagnostic challenge when attempting to differentiate the AH from pathological cardiac adaptation associated with diseases that predispose an athlete to an increased risk of sudden cardiac death (SCD) [8–10].

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In any individual, cardiac adaptation is influenced by a range of training exposure and individual factors [4]. A key mediating variable is the nature of the training exposure (mode, intensity, duration and volume) [4]. To date many studies have attempted to differentiate the cardiac response to endurance versus resistance training [2, 11] with most data supportive of a significantly greater cardiac adaptation to dynamic or endurance based exercise training [2, 7, 11–13] with limited or no significant cardiac adaptation to resistance training [11, 12]. In addition, recent work has demonstrated that dynamic exercise training is the primary driver for cardiac adaptation [11, 13]. The American College of Cardiology Task Force/Mitchell classification [14, 15] categorized sporting disciplines into nine main groups based on their physiological demands with a matrix of 3 levels of dynamic training (low, medium and high) as well as 3 levels of static training (low, medium and high). Given the likely impact of dynamic or endurance training, a clear assessment of RV phenotype in athletes grouped in low, medium and high dynamic training exposure would be valuable to inform the nature of RV remodelling in athletes. Another factor that



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impacts on the interpretation of AH data is the body size and composition of the athletes. To date most AH studies have presented absolute RV structural data or indices scaled linearly to body surface area (BSA) [10]. It is well established that allometric scaling is recommended to provide accurate size-independent measures of cardiac size that will aid accurate interpretation [16, 17].

Technical developments have also had a significant impact on measurement of the AH phenotype. Echocardiography plays an important role in the assessment of RV structure and function. Normal resting RV function has been reported in most athlete studies [10, 11, 13]. Technical advancements such as tissue-Doppler and strain ( $\epsilon$ ) imaging have led to new global and regional indices of RV function and mechanics that may provide extra diagnostic tools in the differentiation of AH from pathological cardiac adaptation [11].  $\epsilon$  imaging has highlighted subtle regional differences in RV mechanics, particularly in those athletes with a more marked RV phenotype [5, 18]. In addition, much of RV  $\epsilon$  data has been reported from the RV lateral wall with limited attention given to septal contribution to RV function.

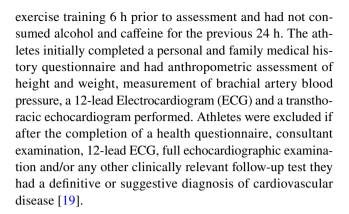
On the basis of these questions and the application of tissue Doppler and  $\varepsilon$  imaging, the aim of this study was to establish the nature and extent of RV structural, functional and mechanical presentation in elite male athletes involved in sports categorised with different levels of dynamic workload in accordance with Mitchell classification [14, 15].

### Methods

## Study design

A prospective cross-sectional study design was performed. Male elite athletes between the age of 18–30 years were included if they were competitive at National level and had no history, signs or symptoms of cardiovascular, metabolic, renal or respiratory disease. No athlete was taking prescribed medication and there was no evidence of a family history of (SCD) or unexplained death under the age of 40 years. The athletes were grouped according to sporting discipline as per Mitchell classification: Low dynamic (LD)—exercise training exposure at < 50% VO<sub>2max</sub> (including cricket, equestrian, motorcycling, martial arts, and weightlifting); Moderate dynamic (MD)—exercise exposure at 50-75% VO<sub>2max</sub> (including rugby); High dynamic (HD)—exercise training exposure at > 75% VO<sub>2max</sub> (including soccer, basketball, ice hockey, boxing, cycling, rowing and triathlon). Ethics approval was obtained by the Ethics Committee of the Liverpool John Moores University and all athletes provided written informed consent.

Athletes attended the echocardiography laboratory for a single visit, ensuring that they had not undertaken any



## **Procedures**

## **Anthropometric assessment**

All athletes were assessed for height and body mass using a standard scale and stadiometer (SECA 764, Birmingham, UK). Body surface area (BSA) was calculated by using a standardised formula [20]. Systolic and diastolic blood pressure were recorded using an automated sphygmomanometer (DINAMAP 300, GE Medical System, Milwaukee, Wisconsin, USA) after 5 min of quiet seated rest.

#### 12-Lead ECG

A standard resting 12-lead ECG (CardioExpress SL6, Spacelabs Healthcare, Washington US) was carried out in accordance with American Heart Association guidelines [19].

# **Echocardiographic assessment**

Echocardiographic examination was performed using a Vivid Q ultrasound machine (GE Healthcare, Horten, Norway) with a 2.5–5 MHz transducer with the subject in the left lateral decubitas position. All acquisitions were made by two experienced sonographers using an echocardiography protocol in accordance with the American Society of Echocardiography [21, 22]. Offline analysis was performed, after storing the images in a raw Digital Imagine and Communications in Medicine format and exported, using commercial available software (EchoPAC V.110.0.2; GE Helathcare, Horten, Norway). All measurements were made in accordance with American Society of Echocardiography guidelines [22] by the same sonographers.

## Conventional 2D, Doppler and tissue Doppler measures

The size of the RV outflow tract was assessed at the proximal (RVOT-PLAX; RVOT1) and distal (RVOT2) levels from the parasternal long and short axis orientations view. RV inflow



dimensions were recorded at the base (RVD1), mid-cavity (RVD2) as well as the length from base to apex (RVD3) from a modified apical four chamber orientation. In the same view, RV area in diastole (RVDA) and systole (RVSA) were obtained by tracing the RV endocardium both in systole and diastole, allowing the calculation of fractional area change (RVFAC). RV wall thickness (RVWT) was measured from a subcostal view at mid wall level. For clinical utility, all structural variables were scaled to individual differences in BSA using a standard linear model. That aside, it is well established that biological systems follow a non-linear relationship and hence all indices were also scaled allometrically according to the laws of geometrically similarity. This data provides size independent values that allow direct comparison across athletes of different body composition and involves scaling linear dimensions to BSA<sup>0.5</sup> and area measurements directly to BSA [16].

RV longitudinal function was determined using M-mode derived tricuspid annular plane systolic excursion (TAPSE) and pulsed wave TDI, at the RV lateral wall tricuspid annulus to obtain peak myocardial velocities in systole (S'), early diastole (E') and late diastole (A'). Pulsed wave Doppler of the RV outflow tract allowed the assessment of RV velocity time interval (RVOT VTI).

## Speckle tracking echocardiography measures

The modified apical four-chamber view was used for the assessment of longitudinal RV lateral wall and septal  $\varepsilon$  and strain rate (SR). To provide optimal endocardial delineation, images were optimised using depth, gain, compression and sector width. Frame rates were set between 80 and 90 frames per second and the focal point was positioned mid cavity to reduce the impact of beam divergence. For offline analysis, pulmonary valve closure (PVC) was obtained from the pulsed wave Doppler signal at the RV outflow tract. A region of interest was placed around the RV basal lateral wall through to basal septum encompassing the mid and apical wall segments. The software automatically tracked the base, mid and apical segments and presented an assessment of tracking quality with segments being excluded if deemed unacceptable. Regional peak and time to peak RV  $\varepsilon$ , peak systolic SR (SRS'), peak early diastolic SR (SRE') and peak late diastolic SR (SRA') were obtained for each of the 6 myocardial segments and a global value was determined as an average of the base, mid and apical wall segments. A base to apex  $\epsilon$  gradient was calculated for both the septum and the lateral wall as the difference between the peak values at both sites.

Temporal  $\epsilon$  and SR was obtained by exporting the raw data to an excel spreadsheet (Excel, Microsoft Corp, Washington, US) and the global temporal values underwent cubic spline interpolation in both systole and diastole phase. The

 $\epsilon$  values were split into 5% increments of the cardiac cycle ensuring the raw peak  $\epsilon$  and SR value was included.

# **Statistical analysis**

Statistical analysis was performed using Statistical Package for the Social Sciences (SPSS) (version 23.0, Chicago IL, USA). All parameters are presented as mean  $\pm$  SD. Normal distribution was confirmed using a Kolmogorov–Smirnov test. Analysis between groups was established using one-way ANOVA and Bonferroni post hoc test for multiple comparison to estimate pairwise differences between groups. Where data was not normally distributed ANOVA on ranks (Krusakal–Wallis) followed by the Mann–Whitney test for pairwise comparisons were employed. To minimize type 1 error rate inflation due to multiple testing, statistical significance was defined as two-sided p < 0.01. Where derived  $\epsilon$  and SR values were significantly different between groups a standard bivariate correlation was used to establish any relationship to absolute RV chamber size.

# **Results**

492 male athletes were recruited (LD: n=48; MD: n=157; HD: n=287) through a range of pre-participation screening events and none were excluded after appropriate follow-up testing. Participant ethnicity was 84% White, 9% Black, 1% Asian, and 6% Mixed. All athletes were matched for training duration and weekly training hours (Table 1). MD athletes had significantly greater weight and BSA compared to both LD and HD groups.

**Table 1** Participant demographics (data are mean  $\pm$  SD)

	LD	MD	HD
Sample (n)	48	157	287
Age (Years)	$25 \pm 3*^{\dagger}$	$23 \pm 4$	$23\pm4$
BSA (m <sup>2</sup> )	$1.9\pm0.2$	$2.1 \pm 0.2^{\ddagger \S}$	$1.9 \pm 0.2$
Height (cm)	$1.8 \pm 0.1$	$1.8 \pm 0.1$	$1.8\pm0.1$
Weight (kg)	$77 \pm 15$	$94 \pm 11^{\ddagger \$}$	$77 \pm 10$
SBP (mmHg)	$131 \pm 8$	$131 \pm 9$	$128\pm12$
DBP (mmHg)	$73 \pm 6*$	$68 \pm 7$	$70\pm8$
Heart Rate (bpm)	$66 \pm 15^{*\dagger}$	$57 \pm 10$	$55 \pm 10$
Training (years)	$10\pm6$	$13 \pm 4$	$13 \pm 6$
Training (hours/week)	$21 \pm 12$	$19\pm8$	$17 \pm 8$

BSA body surface area, SBP systolic blood pressure, DBP diastolic blood pressure

\*P<0.01: LD versus MD

†P<0.01: LD versus HD

‡P<0.01: MD versus LD

§P<0.01: MD versus HD



Absolute and scaled RV structural parameters are presented in Table 2. MD athletes had significantly larger absolute values for RVOT1, RVOT2, RVDA and RVWT than both HD and LD whilst also having larger values of RVOT-PLAX, RVD1 and RVSA when compared to LD alone. HD had significantly larger absolute RVOT2, RVD1, and RVDA compared to LD. Following allometric scaling RVOT-PLAX, RVOT2, RVD1, and RVD3 were larger in HD compared to MD and LD athletes, RVDA was larger in MD and HD compared to LD and RVSA was larger for HD compared with LD athletes.

Standard RV functional data as well as global  $\epsilon$  and SR data are presented Table 3. There were no significant differences between groups for functional and mechanical indices

**Table 2** Absolute and scaled RV structural parameters (data are  $mean \pm SD$ )

	LD	MD	HD
RVOT-PLAX (mm)	$30 \pm 4$	$33 \pm 5^{\ddagger}$	32±4
RVOT1 (mm)	$31 \pm 4$	$35 \pm 5^{\ddagger \$}$	$33 \pm 4$
RVOT2 (mm)	$24 \pm 3$	$26 \pm 4^{\ddagger \$}$	$26 \pm 4^{ll}$
RVD1 (mm)	$41 \pm 5$	$44 \pm 5^{\ddagger}$	$44 \pm 5^{11}$
RVD2 (mm)	$31 \pm 4$	$32 \pm 5$	$31 \pm 6$
RVD3 (mm)	$87 \pm 9$	$90 \pm 11$	$90 \pm 9$
RVDA (cm <sup>2</sup> )	$24 \pm 5$	$29 \pm 4^{\ddagger \$}$	$27 \pm 5^{11}$
RVSA (cm <sup>2</sup> )	$13\pm3$	$16 \pm 3^{\ddagger}$	$15 \pm 3^{11}$
RVWT (mm)	$3.7 \pm 0.8$	$4.1 \pm 0.8^{\ddagger \S}$	$3.9 \pm 0.9$
RVOT-PLAX (mm/m <sup>2</sup> )	$16\pm2$	14 <u>+</u> 4	$16 \pm 2^{9}$
RVOT-PLAX $(mm/(m^2)^{0.5})$	$21 \pm 3$	$22\pm5$	$23 \pm 3^{\text{ll}}$
RVOT1 (mm/m <sup>2</sup> )	$16\pm3$	$15 \pm 4$	$17 \pm 3^{\P}$
RVOT1 $(mm/(m^2)^{0.5})$	$23 \pm 3$	$23 \pm 5$	$24 \pm 3$
RVOT2 (mm/m <sup>2</sup> )	$13\pm2$	$12 \pm 3$	$13 \pm 2^{9}$
RVOT2 $(mm/(m^2)^{0.5})$	$17 \pm 2$	$17 \pm 4$	$19 \pm 3^{ll}$
RVD1 (mm/m <sup>2</sup> )	$21 \pm 3*$	$19 \pm 5$	$22 \pm 3^{\P}$
RVD1 $(mm/(m^2)^{0.5})$	$30\pm3$	$29 \pm 6$	$31 \pm 3^{\text{ll}}$
RVD2 (mm/m <sup>2</sup> )	$16 \pm 2*$	14 <u>+</u> 4	$16 \pm 3^{\P}$
RVD2 $(mm/(m^2)^{0.5})$	$22\pm3$	$21 \pm 5$	$22 \pm 4$
RVD3 (mm/m <sup>2</sup> )	$45 \pm 6*$	$39 \pm 10$	$46\pm5^{\P}$
RVD3 (mm/(m <sup>2</sup> ) <sup>0.5</sup> )	$62 \pm 6$	$59 \pm 12$	$64 \pm 7^{119}$
RVDA $(cm^2/m^2)$	$12\pm2$	$14 \pm 2^{\ddagger}$	$14\pm2^{ll}$
RVSA (cm <sup>2</sup> /m <sup>2</sup> )	7±2	7 ± 1	$8 \pm 2^{ll}$

RV outflow tract diameter at the parasternal long-axis view (RVOT-PLAX), RV outflow tract diameter at the subpulmonary region (RVOT1); at the pulmonic valve annulus (RVOT2), Basal RV diameter (RVD1), Mid RV diameter (RVD2), Base-to-apex length (RVD3), RV end-diastolic area (RVDA), RV end-systolic area (RVSA) and RV wall thickness (RVWT)

\*P<0.01: LD versus MD  $^{\ddagger}$ P<0.01: MD versus LD  $^{\S}$ P<0.01: MD versus HD  $^{\P}$ P<0.01: HD versus MD  $^{\parallel}$ P<0.01: HD versus LD



**Table 3** RV functional parameters (Data are mean  $\pm$  SD)

	LD	MD	HD
TAPSE (mm)	23±5	24±4	24±4
RVFAC (%)	$46 \pm 8$	$46 \pm 8$	$45\pm8$
RVOT VTI (cm)	$19\pm3$	$18 \pm 3$	$20 \pm 16$
TDI S' (cm/s)	$15\pm3$	$15\pm2$	$14\pm2$
TDI E' (cm/s)	$15\pm3$	$15 \pm 3$	$15 \pm 3$
TDI A' (cm/s)	$10\pm3$	$10\pm3$	$10\pm3$
Time to peak S' (ms)	$183 \pm 21$	$193 \pm 23$	$190 \pm 29$
Time to Peak E' (ms)	$509 \pm 49$	$526 \pm 45$	$522 \pm 54$
Peak ε (%)	$-23.4 \pm 3.1$	$-22.7 \pm 2.7$	$-23.5 \pm 2.6$
Time to Peak $\varepsilon$ (s)	$0.37 \pm 0.04$	$0.38 \pm 0.03$	$0.38 \pm 0.04$
Peak SRS' (l/s)	$-1.2 \pm 0.2$	$-1.1 \pm 0.2$	$-1.2 \pm 0.2$
Peak SRE' (l/s)	$1.6 \pm 0.3$	$1.5 \pm 0.3$	$1.6 \pm 0.3$
Peak SRA' (l/s)	$0.7 \pm 0.2$	$0.8 \pm 0.2$	$0.7 \pm 0.2$

Tricuspid Annular Plane Systolic Excursion (TAPSE), RV Fractional Area Change (RVFAC), RV Outflow Tract (RVOT), RV outflow Tract Velocity Time Integral (RVOT VTI), Peak systolic motion on Tissue Doppler Image (TDI RV S'), early diastolic motion (TDI RV E'), late diastolic motion (TDI RV A'), Strain Rate during ventricular early diastole (SRE'), Strain Rate during late ventricular late diastole (SRA') and Strain Rate during ventricular systole (SRS')

derived from standard conventional 2D, tissue Doppler and  $\epsilon$  imaging. Segmental mechanical analysis determined that MD athletes had significantly lower  $\epsilon$  in the apical septal wall segment compared to both the HD and LD groups. This contributed to a significantly lower base-apex  $\epsilon$  gradient in the septal wall of MD athletes (Table 4). Assessment of the temporal distribution of global  $\epsilon$  and SR across the cardiac cycle shows no significant differences between the three groups and is presented in Fig. 1.

**Table 4** RV strain ( $\epsilon$ ) base to apex gradient (data are mean  $\pm$  SD)

	LD	MD	HD
Global	$-23.4 \pm 3.1$	$-22 \pm 5.2$	$-22.6 \pm 5$
Base to apex gradient ε (%) (lateral Wall)	$3.9 \pm 8.5$	$2.8 \pm 7.8$	$5.7 \pm 7.5$
Basal	$-26.4 \pm 6.6$	$-26.4 \pm 7.1$	$-25.9 \pm 5.5$
Mid	$-27.3 \pm 4.8$	$-27.3 \pm 5.4$	$-27.8 \pm 4.2$
Apical	$-30.9 \pm 4.5$	$-28.7 \pm 5$	$-31.4 \pm 4.9^{\P}$
Base to apex gradient ε (%) (septal Wall)	$-5.6 \pm 5.3$ *	$-0.4 \pm 5.5$	$-3.2 \pm 5.8^{\P}$
Apical	$-22.9 \pm 4.5*$	$-18.8 \pm 4.8$	$-21.5 \pm 4.7^{\P}$
Mid	$-18.3 \pm 3.1$	$-17.8 \pm 2.5$	$-17.6 \pm 2.5$
Basal	$-17.6 \pm 2.7$	$-18.6 \pm 2.7$	$-18\pm3.1$

\*P<0.01: LD versus MD ¶P<0.01: HD versus MD

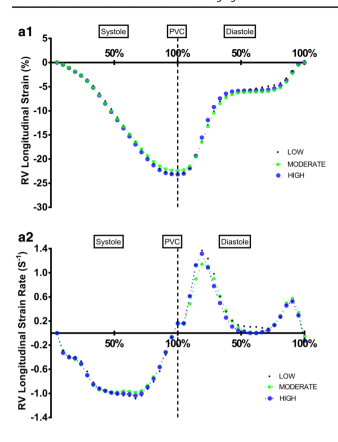


Fig. 1 Right ventricular temporal curves of a1 mean longitudinal  $\epsilon$  and a2 strain rate compared between Low, Moderate and High dynamic classifications

There were significant positive, but small correlations between the apical septal  $\epsilon$  and RVOT-PLAX (r=0.14, P=0.014), RVOT1 (r=0.11, P=0.041), RVOT2 (r=0.13, P=0.015) and RVDA (r=0.13, P=0.016). Apical lateral  $\epsilon$  was correlated with RVOT2 (r=0.12, P=0.03).

## **Discussion**

The key findings from this study were (1) athletes involved in sports with MD and HD training exposure had generally larger absolute and scaled RV structural indices than athletes with LD training exposure, (2) there were no between differences in conventional RV functional indices as well as global RV  $\varepsilon$  and SR, (3) athletes with MD training exposure had a smaller base to apex  $\varepsilon$  gradient in both septal and lateral walls largely due to reduced apical  $\varepsilon$  and this appears to be partly related to absolute RV chamber size.

# Right ventricular structure

It has been established that exercise training in elite male athletes causes RV structural adaptation [2, 6, 9, 23–26], which is likely driven by changes in cardiac load [27].

During acute dynamic exercise the body employs large muscle groups that require substantial O<sub>2</sub> demand. This demand is met by increased cardiac work elevating both LV and RV cardiac output. The long-term adaptation to this acute loading is a significant dilation of the ventricular cavities and this is clearly observed in the RV of the HD and MD athletes in this study. This RV data supports a range of previous research in largely endurance-based athletes [2, 13, 26, 28] as well as data from a training study Spence et al. [28] that identified that participants engaged in a HD exercise training programme had greater RV adaptation than those with LD exercise training. A recent meta-analysis partially supports our work with strength trained athletes (LD) having smaller RV inflow chamber size [26]. Interestingly, unlike our data, this meta-analysis demonstrates no training specific differences in the RV outflow. The absolute RV structural data in this study were generally greater in the MD athletes, which may be somewhat counterintuitive based on training loads. It is important to note, however, that the MD athletes were physically bigger and that after scaling both HD and MD athletes had larger RV structural indices than the LD athletes. Utomi et al. [11] undertook a meta-analysis detailing greater RV enlargement in endurance athletes compared to control participants, however, there was a positive and significant association in the regression model between RV parameters and BSA. Our data confirm the importance and utility of appropriate scaling of cardiac data [17]. It is interesting to note that many of the MD athletes were elite rugby football league players. As well as being bigger athletes, it may be pertinent to question whether the American College of Cardiology/Mitchell Classification have placed this athletic group in the wrong category. It is pertinent to recommend that all RV structural indices in the athlete should be indexed appropriately to provide size independence.

### Right ventricular function

Our conventional echocardiographic data demonstrated no difference in global RV function, during systole or diastole, between groups undertaking different amounts of dynamic training. Although there is evidence to suggest athletes have different RV function than non-athlete controls [2, 5, 29–32], there are limited data between different athlete groups. Those studies that have compared athletes largely report no difference in TAPSE [2, 13, 26, 33], RVFAC [2, 25, 34] and myocardial tissue velocities [2, 7, 34]. Similar to these conventional indices of global RV function there was no difference in peak global  $\epsilon$  and SR between athlete groups which supports some previous studies [2, 13, 26].

There is less data in the literature reporting regional RV  $\varepsilon$  and SR in dynamically trained athletes. The observation of reduced apical  $\varepsilon$  in the RV septal wall segment in MD athletes is a novel finding and appears to be related, in part,



to RV size and potentially body size. Teske et al. [5] and La Gerche et al. [18] both demonstrated reduced RV basal SR in athletes with larger RV area and attributed this to RV geometry, heart rate and resting reserve [5, 18]. Teske et al. [35] reported a reduction in  $\varepsilon$  and SR in basal and mid segments in endurance athletes with marked RV dilatation as normal physiological adaptation [35]. In addition, during the competitive season, D'Ascenzi et al. [36] documented no significant differences (P > 0.14) in global  $\varepsilon$  among 29 toplevel athletes. They did however demonstrate an increase in RV apical wall  $\varepsilon$  at both mid-season and end-season which was associated to RV mid-cavity, RVEDA, RVESA and SV [36]. The impact of hand grip exercise on RV apical  $\varepsilon$  has also been demonstrated in an acute setting and may provide some evidence for this type of stimulus leading to chronic functional adaptation [37]. During maximal exercise the RV has the ability to increase its contractility to compensate for the increase in work [38] and it is likely that the changes in the RV apical segment  $\varepsilon$  is due to the changes in training intensity and chamber size. The data from the current study demonstrates an association between apical septal  $\epsilon$ and RV outflow tract, rather than the inflow/sinus region of the RV. This might provide some explanation for the reduction in apical deformation, rather than that of basal segments between different dynamic sport discipline. Evidence shows crisscross fibres within the apical interventicular septum which contribute to the RVOT structure [39]. Therefore, any structural changes in the RVOT might contribute to the apical  $\varepsilon$  in this wall segment. It is clear that RV size accounts for only a small portion of the variance in the apical septal  $\varepsilon$ and thus further work is required to provide further mechanistic insight.

# Clinical/imaging implications

It is documented that 3–10% of all sudden cardiac deaths in young athletes is due to arrhythmogenic right ventricular cardiomyopathy (ARVC) [40]. In this regard, differentiation of RV physiological adaptation from pathological changes caused by ARVC is of utmost importance in pre-participation cardiac screening programs. This study provides valuable information on RV structural and functional adaptation in athletes undertaking different amounts of dynamic load exercise training, which may aid in specific screening programs. This study presents absolute and scaled RV structure that shows the importance of indexing to provide size-independent RV structural data.

These data demonstrate that despite marked RV dimensional remodelling, exercise-induced RV adaptation in a young male elite athlete population is not associated with any degradation in function and thus does not support the theory of an exercise-induced ARVC. It is important to note that any potential mal-adaptation occurs in sporadic

cases and may be dependent on long term repeated exposure and insufficient recovery time [41].

The addition of novel non-invasive imaging techniques may provide unique insights into the RV AH phenotype particularly the potential use of speckle-tracking echocardiography [42, 43]. Our data also demonstrates the use of both septal and free wall  $\epsilon$  for assessment of global RV function. This is challenging due to the inability to clearly define which portion of the septum contributes to the RV. It is therefore apparent that further work in athletic and non-athletic populations should aim to establish the extend and contribution of septal and free wall deformation to global RV mechanics.

## Limitations

This study explored the effect of three different levels of dynamic exercise load upon the RV without taking into account the static components for each level (nine box Mitchell classification), which would help determine the effect of varying static and dynamic components of each sport on cardiac adaptation. There is some discrepancy in sample size in the 3 groups but larger samples in all groups would have improved the power and generalisability of the data. The addition of a sedentary control group would also provide some benefit. Furthermore, this study only investigated RV remodelling among male athletes It is apparent that further study related to gender specific adaptation would be valuable.

# **Conclusion**

This study compared the effects of 3 different levels of dynamic exercise training (using the Mitchell classification) on RV structure, function and mechanics. RV morphology was generally larger in HD athletes, compared to MD and LD groups when data were allometrically scaled to BSA. Despite no significant differences in RV function between the three groups, there was reduction in regional  $\epsilon$  in the apical septal wall segment which may partially be explained by greater absolute RV dimensions and body size in MD athletes.

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#### Compliance with ethical standards

**Conflict of interest** There is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.



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