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**Kleinnibbelink, G, Panhuyzen-Goedkoop, N, Hulshof, HG, van Dijk, A, George, KP, Somauroo, J, Oxborough, D and Thijssen, DHJ**

**Exercise Training Induces Left- but not Right-sided Cardiac Remodelling in Olympic Rowers**

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### Article

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1 **Exercise Training Induces Left- but not Right-Sided Cardiac Remodelling in Olympic**  
2 **Rowers**

3 **Running title: Cardiac remodelling in Olympic rowers**

4

5 **ABSTRACT**

6 Whilst the athlete's heart has been extensively described, less work has focused on the potential  
7 for elite athletes to demonstrate further cardiac remodelling upon an increase in training  
8 volume. Moreover, little work explored potential side-specific cardiac remodelling. Therefore,  
9 we examined the impact of an increase in training volume across 9-months in elite rowers on  
10 left- and right-sided cardiac structure, function and mechanics (i.e. longitudinal, radial and  
11 circumferential strain, twist and strain-volume loops). As part of the preparations to the 2012  
12 Olympic Games, twenty-seven elite rowers ( $26.4\pm 3.7$  years, 19 male) underwent  
13 echocardiography prior to and post (9-months) an increase in training volume (24 to 30-35h  
14 weekly). Training increased left ventricular LV structure, including wall thickness, diameter,  
15 volume, mass and LV twist (all  $p<0.05$ ). Female rowers demonstrated larger adaptation in left  
16 ventricular diameter and mass compared to male rowers (both  $p<0.05$ ). No changes were  
17 observed in other measures of left ventricular function in both sexes (all  $p>0.05$ ). The 9-month  
18 intervention showed no change in right ventricular/atrial structure, function or mechanics (all  
19  $p>0.05$ ). In conclusion, our data revealed that 9-month increased training volume in elite  
20 rowers induced left-sided (but not right-sided) structural remodelling, concomitant with an  
21 increase in left ventricular twist, with some changes larger in women.

22 **Keywords:** athlete's heart; right ventricle; cardiovascular disease; echocardiography; speckle  
23 tracking echocardiography

## 24 INTRODUCTION

25 Exercise training represents a potent stimulus for remodelling of the heart. Recent prospective  
26 and long-term intervention studies support the presence of predominant eccentric ventricular  
27 adaptation in response to exercise training [1, 2]. In other words, regular exercise training leads  
28 to a balanced increase in both volume and mass, whilst function seems largely preserved [1].  
29 Previous work largely focused on either adaptation of the left side or adaptation of the right  
30 side of the heart in response to exercise training. As such, there is an important gap in the  
31 literature pertaining to the lack of knowledge whether exercise training differentially affects  
32 the left *versus* right ventricle in elite athletes.

33 Both ventricles receive a similar amount of blood. Due to their distinct geometry and mass,  
34 with the right ventricle (RV) being larger in volume but smaller in wall thickness than the left  
35 ventricle (LV), both ventricles may be exposed to distinct hemodynamic stimuli, potentially  
36 leading to different patterns of adaptation in both structure and function [3]. Indeed, exercise  
37 leads to distinctly different changes in afterload for both ventricles, with relatively larger  
38 increases in afterload for the RV [4]. This supports the potential for different adaptation  
39 between ventricles. Better insight into differences in remodelling between ventricles is highly  
40 relevant, especially since previous studies have linked exercise-induced RV cardiomyopathy  
41 to high volumes of exercise training in elite athletes [5, 6]. Insight into the potential presence  
42 of side-specific physiological remodelling of the heart will also contribute to improved  
43 interpretation of pre-participation screening for high-risk cardiovascular conditions associated  
44 with sudden cardiac arrest in athletes.

45 The aim of this study, therefore, was to examine the impact of an increase in volume (across  
46 9-months) in elite rowers on left- and right-sided cardiac structure, function and mechanics (i.e.  
47 longitudinal, radial and circumferential strain, twist and strain-volume/area loops). Based on

48 the higher relative workload for the RV [4], we expect larger structural cardiac adaptation in  
49 the RV (whilst preserving function) compared to the LV in elite rowers.

50 Previous work suggested that sex may differentially affect cardiovascular function during  
51 physiological stimuli [8, 9]. Based on cross-sectional comparisons, similar patterns of cardiac  
52 remodelling have been observed with static and mixed exercise between men and women,  
53 whilst greater LV adaptation may be present in women during dynamic exercise [7, 10].  
54 Therefore, as an explorative aim, we evaluate the impact of sex on the impact of exercise in  
55 elite rowers on left- and right-sided cardiac structure, function and mechanics

56

## 57 **METHODS**

### 58 **Study population and study design**

59 In this prospective, longitudinal study, as part of the work-up to the 2012 Olympic Games,  
60 twenty-seven elite level rowers (male and female, all Caucasian) underwent baseline  
61 echocardiography prior to and post (9-months) a planned increase in training volume. Baseline  
62 echocardiograms were performed immediately after the 2011 World Rowing Championships  
63 (i.e. when all athletes were in a highly trained status), and 3 months before the 2012 Olympic  
64 Games. After the baseline echocardiograms, the rowers, both male and female, increased their  
65 training volume gradually from 24 hours to 30-35 hours per week (20% strength, 80% rowing  
66 training consisting of high-intensity interval and endurance training). Height and weight were  
67 obtained before echocardiography was performed (SECA scale and stadiometer, SECA GmbH,  
68 Hamburg, Germany). This study was conducted in accordance to with the ethical standards in  
69 sport and exercise science research and approved by the [REDACTED]  
70 ethics committee [11].

71

## 72 **Echocardiographic measurements**

73 The echocardiographic examinations were performed in the left lateral decubitus position by  
74 one highly experienced cardiologist (AvD) using a Vivid-Q ultrasound machine (GE Medical,  
75 Horton, Norway), equipped with a 1.5-4 MHz phased array transducer. Heart rate was  
76 calculated from a single lead ECG inherent to the ultrasound system. Images were stored in  
77 raw digital imaging and communication in medicine (DICOM) format and were exported to an  
78 offline workstation (EchoPac, version 113, GE Medical, Horton, Norway). Data-analysis, from  
79 three stored cycles, was performed by a single observer with experience in echocardiography  
80 (GK) using commercially available software (EchoPac, version 113, GE Medical, Horton,  
81 Norway). The echocardiograms were all coded so the observer was blinded for the timing (pre  
82 vs. post) and for sex (male vs. female).

83 *Conventional measurements.* Cardiac structural and functional measurements were made  
84 according to the current guidelines for cardiac chamber quantification [12]. Regarding the left  
85 heart, we examined the following structural and functional indices: wall thickness of the  
86 septum (IVSd) and posterior wall (PWd), internal cavity diameter at end-diastole (LVIDd), LV  
87 mass (LVM), anteroposterior diameter of the left atrium (LA), LA volume by the disk  
88 summations technique in apical 4-chamber (A4C) and apical 2-chamber (A2C) view, modified  
89 Simpson's left ventricular ejection fraction (LVEF), tissue Doppler imaging (TDI) of the mitral  
90 annulus ( $s'$ ,  $e'$  and  $a'$ ) and trans-mitral Doppler (E, A and E/A ratio). Regarding the right heart,  
91 following structural and functional indices were determined: basal and mid-cavity end-diastolic  
92 diameters, RV end-diastolic area (RVEDA), RV end-systolic area (RVESA), RV outflow tract  
93 (RVOT) diameter at the proximal level in the parasternal long-axis (PLAX) and parasternal  
94 short-axis (PSAX) view, right atrial (RA) area, RV fractional area change (RVFAC), tricuspid  
95 annular plane systolic excursion (TAPSE), TDI of the tricuspid annulus. All LV and RV

96 structural indices were allometrically scaled to body surface area (BSA) according to the laws  
97 of geometric similarity [13].

98 *Mechanics.* Images were acquired specifically for speckle tracking. This involved the  
99 optimization of frame rates between 40 and 90 frames s<sup>-1</sup>, depth to ensure adequate imaging of  
100 the chamber of interest, a focal zone at mid-cavity to reduce the impact of beam divergence  
101 and gain, compression and reject to ensure endocardial delineation.

102 *Ventricular and atrial mechanics.* The A4C view was utilized for LV, LA and RA global  
103 longitudinal strain and the RV focused view for the RV longitudinal strain. The LV short-axis  
104 (SAX) views (basal, mid and apical) were utilized for radial, circumferential strain and twist.  
105 Valve closure times were determined from the respective pulsed wave Doppler signals.

106 For all compartments (LV, LA, RV, RA), the myocardium was manually traced and adjusted  
107 so that the region of interest (ROI) incorporated all of the wall thickness, while avoiding the  
108 pericardium. The region of interest was divided into six myocardial segments, providing  
109 segmental strain curves and a global longitudinal strain curve as an average of all six segments.  
110 In order to obtain peak LV circumferential strain, peak LV radial strain and peak apical and  
111 basal rotation, a full-thickness ROI of the mid-, basal- and apical-SAX views, which was  
112 divided into six segments, was selected. In addition, raw strain values were exported and a  
113 cubic spline was applied to normalize for heart rate. This allowed the presentation of temporal  
114 strain and rotation across the cardiac cycle.

115 *Strain-volume/area loops.* The longitudinal strain-volume/area relationship (for methodology  
116 of derivation, see Supplemental 1 and Oxborough *et al.* [14]) was assessed using the following  
117 parameters: (I) the linear strain-area slope (Sslope) and early strain-area slope during first 5%  
118 of volume ejection in systole (ESslope); (II) end-systolic peak longitudinal strain (peak strain);  
119 (III) the early linear strain-area slope during first 5% (EDslope) and late linear strain-area slope

120 (LDslope) during last 5% of volume increase in diastole; and (IV) diastolic uncoupling (i.e.  
121 difference in strain between systole and diastole at any given area), divided into uncoupling  
122 during early (Uncoupling ED) and late diastole (Uncoupling LD) [14, 15].

123

## 124 **Statistical analysis**

125 Statistical analyses were performed using SPSS Statistics 24 (SPSS, Inc., Chicago, Illinois).

126 All parameters were visually inspected for normality and tested with Shapiro-Wilk normality  
127 tests. Continuous variables were reported as mean±SD and categorical variables were  
128 presented as proportions. Paired-Samples T-tests were used to compare echocardiographic  
129 continuous variables between the baseline and follow-up evaluation. Comparison of sex  
130 differences was performed using repeated measurements ANOVA with Bonferroni *post hoc*  
131 correction for multiple comparisons.

132 Consistency of intra-observer measurements of selected measurements were verified through  
133 the intra-class correlation coefficient (ICC). Therefore, both echocardiographs of 15 randomly  
134 chosen subjects were analysed by the same operator blinded from earlier results. ICC  
135 coefficients were as follows: RVOT-PLAX 0.96 (0.91-0.98), LVISd 0.91 (0.81-0.96), LVPWd  
136 0.90 (0.80-0.95), LVIDd 0.97 (0.93-0.98), RV basal diameter 0.98 (0.96-0.99), RVEDA 0.98  
137 (0.95-0.99), RA area 0.99 (0.97-0.99), LA volume 0.98 (0.95-0.99). In previous studies, we  
138 showed that strain measurements and both right and left ventricular loops have a good to  
139 excellent inter-observer variability [14-17].

140

## 141 **RESULTS**

### 142 **Baseline characteristics**



143 All 27 rowers participated in the 2012 Olympic games. Mean age of the study population was  
144  $26.4 \pm 3.7$  years, consisting of 19 males (70%,  $26.3 \pm 4.3$ y) and eight females (30%,  $26.6 \pm 1.9$ y).  
145 All rowers were Caucasian. Male rowers were significantly taller ( $193.7 \pm 6.6$  versus  $181.5 \pm 8.8$   
146 cm,  $p=0.001$ ), heavier ( $88.0 \pm 12.0$  versus  $72.9 \pm 8.6$  kg,  $p=0.003$ ) and had greater BSA ( $2.2 \pm 0.2$   
147 versus  $1.9 \pm 0.2$  m<sup>2</sup>,  $p=0.002$ ), but had a similar BMI ( $23.3 \pm 2.1$  versus  $22.2 \pm 1.0$ ,  $p=0.08$ )  
148 compared to female rowers. Weight, body surface area (BSA) and body mass index (BMI) did  
149 not significantly change over time ( $83.9 \pm 13.0$  to  $84.3 \pm 13.0$  kg,  $p=0.10$ ;  $2.1 \pm 0.2$  to  $2.1 \pm 0.2$  m<sup>2</sup>,  
150  $p=0.10$ ;  $23.0 \pm 1.9$  to  $23.1 \pm 1.9$  kg/m<sup>2</sup>,  $p=0.11$ , respectively). Resting heart rate was higher at  
151 follow-up compared to baseline  $54 \pm 7$  to  $58 \pm 8$  bpm ( $p=0.02$ ).

152

### 153 **Exercise training and cardiac remodelling: comparison between sides**

154 *Left ventricle and atrium.* There was a significant increase in LV wall thickness, diameter,  
155 volume and mass (all  $p<0.01$ ), which remained significant after correction for BSA (all  $p<0.05$ )  
156 (**Table 1, Figure 1**). Similarly, there was a significant increase in LA diameter and volume  
157 (both  $p<0.01$ ), which remained significant after correction for BSA (both  $p<0.01$ ). Exercise  
158 training increased LV twist, whilst no other changes in functional or mechanical indices were  
159 found (**Table 1, Figure 1-2**).

160 *Right ventricle and atrium.* We found no significant changes in right ventricular and atrial  
161 structure, function and mechanics (**Table 2, Figure 1-2**).

162

### 163 **Exercise training and cardiac remodelling: comparison between sexes**

164 *Baseline characteristics.* At baseline, female rowers had smaller LV and RV cardiac  
165 dimensions compared to male rowers (all  $p<0.05$ , **Table 1-2**), which was not present after

166 correcting for BSA (all  $p > 0.05$ ). Absolute RVOT dimensions did not differ between sexes  
167 (**Table 2**). Female rowers had a smaller LV mass compared to male rowers ( $p < 0.01$ ), which  
168 remained significant after correction for BSA ( $p < 0.05$ , **Table 1**). Except for a lower TAPSE  
169 and a higher E velocity in female rowers (both  $p = 0.02$ ), no significant differences were found  
170 in conventional measurements of left- or right-sided cardiac function (**Table 1-2**). Female  
171 rowers demonstrated significantly higher LV apical circumferential strain, lower peak systolic  
172 apical rotation (both  $p < 0.05$ ) and steeper slopes of the left- and right-sided strain-volume/area  
173 loop compared to male rowers (LV – Sslope, LDslope, both  $p < 0.05$ ; RV – Sslope, ESlope,  
174 LDslope, all  $p < 0.05$ ) (**Table 1-2, Figure 2**).

175 *Training-induced remodelling.* Females demonstrated a significantly larger increase in  
176 absolute and scaled LV diameter and LV mass compared with male rowers (**Table 1, Figure**  
177 **3**). No differences were found between sexes in the right ventricle or atrium (**Table 2**).

178

## 179 **DISCUSSION**

180 The aim of our study was to examine the impact of an increase in volume (across 9-months) in  
181 elite rowers on left- and right-sided cardiac structure, function and mechanics. We present the  
182 following findings. First, an increased training volume in elite rowers across 9-months resulted  
183 in significant structural adaptation of the left ventricle and atrium, with no adaptations observed  
184 on the right side. Second, left-sided structural cardiac adaptation was accompanied by an  
185 increase in LV twist, but no other left- or right-sided functional adaptations. This highlights the  
186 plasticity of the heart for remodelling in response to exercise training, even in elite athletes.  
187 Taken together, our results demonstrate cardiac side-, and possibly also sex-specific adaptation,  
188 which is relevant for future studies that should acknowledge that cardiac remodelling does not  
189 simply follow the same path between and within individuals.

190

191 After an increase in training volume across 9-months, the left heart of this cohort of elite rowers  
192 showed mild structural (eccentric) adaptation with an increased LV twist, whilst there was no  
193 significant remodelling in the right heart. This left-sided structural adaptation is in line with  
194 several previous longitudinal training studies including sedentary, moderately- and highly-  
195 trained individuals [2, 18-21]. Interestingly, concomitant to left-sided structural adaptation,  
196 elite rowers also demonstrated augmented LV twist after the increase in training volume. The  
197 higher heart rate post-training may partially explain the increase in twist. However, this seems  
198 unlikely since no correlation was found between heart rate and twist ( $r=0.02$ ,  $p=0.94$ ).  
199 Moreover, other functional parameters (also susceptible for differences in heart rate) did not  
200 change over time. Although we found adaptation in functional and structural characteristics,  
201 both may demonstrate a distinct pattern and are not similarly present. Indeed, the increase in  
202 twist was not correlated with changes in LV cardiac morphology (data not shown). Moreover,  
203 *Weiner et al.* observed that exercise training may initially (i.e. 90 days) lead to increases in LV  
204 twist, which subsequently disappeared during the chronic training phase (i.e. 39 months) [22].  
205 Our finding provides some support for this concept, in that an increase in volume of exercise  
206 initially resulted in both functional and structural adaptations, where functional changes may  
207 ultimately normalize during the chronic phase when volume of exercise remains the same.  
208 Future work is required to better understand these time-dependent adaptations in cardiac  
209 remodelling.

210

211 Despite the disproportionate load on the right *versus* left ventricle during exercise [4], we found  
212 no adaptation of the right ventricle or atrium. This finding contrasts with our hypothesis, but  
213 also with others who addressed right-sided cardiac remodelling in elite athletes [23, 24].

214 D'Ascenzi *et al.* reported seasonal variation in RV size in a cohort of top-level basketball and  
215 volleyball players [23]. Across three consecutive Olympic Games, Aengevaeren *et al.* noted  
216 that RV remodelling occurred between the first two Olympics Games, followed by a plateau  
217 during the subsequent 4 years in a heterogeneous group of athletes (n=50, 17 different sports)  
218 [24]. These studies, however, are limited by the impact of ageing (i.e. 8-year cycle), large  
219 variations in training status across the season, and/or the heterogeneous group of athletes  
220 included. The lack of RV remodelling in our study may be explained by achieving a  
221 physiological limit for further adaptation prior to the start of the increase in training volume in  
222 highly trained rowers due to pericardial constraint. At least, our observations support the  
223 presence of distinct remodelling between the left and right side of the athlete's heart. Future  
224 work is required to better understand these differences, specifically focusing on the distinct  
225 load placed on both ventricles during exercise, possibly underlying distinct remodelling to  
226 (high) volumes of exercise training.

227

228 Following the explorative analysis, this study examined the impact of sex on cardiac adaptation  
229 to training using a longitudinal design. This design markedly differs from most previous studies  
230 that have adopted a cross-sectional design, including a heterogeneous groups of athletes, and  
231 generally not using allometric scaling [10, 25-29]. Our data showed larger LV structural  
232 adaptation in female rowers, which remained present upon allometric scaling. These distinct  
233 adaptations cannot relate to differences in lifetime exposure to elite athlete level training, since  
234 both groups do not differ in age (males  $26.3 \pm 4.3$ y *versus* females  $26.6 \pm 1.9$ y,  $p=0.84$ ).  
235 Alternative explanations for the distinct remodelling might be hormonal, molecular and/or  
236 genetic mechanisms. However, these mechanisms are not fully understood yet and represent  
237 topics for future research [7]. Another explanation is the potential for differences in the  
238 (absolute and relative) workload of the exercise training between men and women.

239 Unfortunately, data were not available to compare exercise intensity and workload between  
240 groups or individuals. An important limitation is the relatively low sample size for female  
241 rowers within this analysis. Nonetheless, our study was sufficiently powered to detect a  
242 significant effect between sexes in adaptation. We performed *post hoc* calculations and found  
243 that our study has a statistical power of 0.51-0.64 to detect sex differences in LV mass and LV  
244 diameter. At least, our findings highlight the importance for future research to better understand  
245 and establish potential sex differences in cardiac adaptation in response to exercise training.

246

247 *Clinical relevance.* The observation of no further adjustment in RV remodelling seems relevant  
248 as RV enlargement may overlap with the pathological dilation of the RV in patients with an  
249 arrhythmogenic RV cardiomyopathy. Previous studies have related RV remodelling in the  
250 already highly trained athlete to potential clinical problems. Our work suggests that even high  
251 volumes of exercise does not automatically lead to further remodelling of the RV, despite  
252 structural changes in the LV. However, a potential limitation is that subjects followed an  
253 individually determined exercise training protocol to increase training volume, which makes it  
254 difficult to relate cardiac remodelling to specific determinants of the exercise training protocol  
255 and at a cohort level. However, all individuals significantly increased their training volume,  
256 highlighting that additional cardiac remodelling is possible upon increases in training volume.  
257 Our study may have further clinical relevance, since we specifically explored remodelling in  
258 female elite athletes. Participation of females in elite sports has increased significantly over the  
259 past decades. Current work on the athlete's heart, leading to insight into (ab)normal levels of  
260 adaptation, largely originate from studies performed in males. Our work supports performing  
261 specific studies in women, examining the geometry and potential pathological relevance of the  
262 female athlete's heart.

263

264 In conclusion, our data suggest that an increased exercise training volume in elite rowers across  
265 9-months induced side-specific cardiac remodelling. Specifically, we found left-sided (but not  
266 right-sided) structural adaptations, with concomitant increase in LV twist in already highly  
267 trained rowers. Interestingly, these adaptations were significantly larger in women compared  
268 to men, a finding that warrants further exploration in future work. Taken together, our work  
269 suggests that examining the athlete's heart should go beyond the single-sided approach most  
270 previous studies adopted, and should explore both left and right-sided adaptation.

271

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275 None

## 276 **CONFLICT OF INTEREST/DISCLOSURES**

277 The authors have no relationships or conflicts to disclose.

278

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**TABLES LEGENDS**

**Table 1 – Left heart structural, functional and mechanical echocardiographic parameters observed in male and female elite rowers prior to and post a 9-month increase in training volume**

**Table 2 – Right heart structural, functional and mechanical echocardiographic parameters observed in male and female elite rowers prior to and post a 9-month increase in training volume**

## FIGURES LEGENDS

### **Figure 1 – Structural and functional cardiac remodelling after 9-months training volume increase**

Figure 1: Change in right (A-C-E) and left (B-D-F) sided cardiac structure and function in elite rowers (n=27) before ('Pre': black bars) and after ('Post': grey bars) a 9-month training period. Error bars represent SD. \* significantly different from pre (p<0.05).

### **Figure 2 – Strain-Area/Volume loops**

Figure 2A: Left and right ventricular strain-volume/area loop in elite rowers before ('Pre Systolic': black lines, 'Pre Diastolic': black dotted lines) and after ('Post Systolic': red lines, 'Post Diastolic': red dotted lines) a 9-month training period. 2B: Strain-volume/area loops distributed to sex.

### **Figure 3 – Cardiac remodelling distributed to sex**

Figure 3: Change in (A) LV mass index and (B) LV diameter index in elite rowers before ('Pre': black bars) and after ('Post': grey bars) a 9-month training period distributed to sex. Error bars represent SD.