Do big athletes have big hearts? Impact of extreme anthropometry upon cardiac hypertrophy in professional male athletes

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ABSTRACT

Aim Differentiating physiological cardiac hypertrophy from pathology is challenging when the athlete presents with extreme anthropometry. While upper normal limits exist for maximal left ventricular (LV) wall thickness (14 mm) and LV internal diameter in diastole (LVIDd, 65 mm), it is unknown if these limits are applicable to athletes with a body surface area (BSA) >2.3 m².

Purpose To investigate cardiac structure in professional male athletes with a BSA>2.3 m², and to assess the validity of established upper normal limits for physiological cardiac hypertrophy.

Methods 836 asymptomatic athletes without a family history of sudden death underwent ECG and echocardiographic screening. Athletes were grouped according to BSA (Group 1, BSA>2.3 m², n = 100; Group 2, 2–2.29 m², n = 244; Group 3, <1.99 m², n = 492).

Results There was strong linear relationship between BSA and LV dimensions; yet no athlete with a normal ECG presented a maximal wall thickness and LVIDd greater than 13 and 65 mm, respectively. In Group 3 athletes, Black African ethnicity was associated with larger cardiac dimensions than either Caucasian or West Asian ethnicity. Three athletes were diagnosed with a cardiomyopathy (0.4% prevalence); with two athletes presenting a maximal wall thickness >13 mm, but in combination with an abnormal ECG suspicious of an inherited cardiac disease.

Conclusion Regardless of extreme anthropometry, established upper limits for physiological cardiac hypertrophy of 14 mm for maximal wall thickness and 65 mm for LVIDd are clinically appropriate for all athletes. However, the abnormal ECG is key to diagnosis and guides follow-up, particularly when cardiac dimensions are within accepted limits.

INTRODUCTION

Regular and prolonged intensive exercise is associated with cardiac morphological adaptation, together with electrocardiographic alterations. Significant cardiac enlargement may be an expression of underlying cardiac disease, placing the athlete at a greater risk of sudden cardiac death (SCD). In rare cases, the degree of physiological adaptation in cardiac morphology can mimic that of a number of pathological disease states, most notably hypertrophic cardiomyopathy (HCM). Differentiation between a physiological or pathological remodelling process is, therefore, extremely important. Consequently, establishing the upper normal limits of physiological enlargement in response to physical training is an important focus for clinicians and scientists.

Fourteen and 65 mm have been established as the physiological upper limits for maximal wall thickness and left ventricular (LV) internal diameter during diastole, respectively. These limits come from three large-scale studies examining approximately 4800 elite athletes (predominantly male), who observed that a small minority (1.5–4%) demonstrated LV hypertrophy (LVH) >13 mm and 4–6% have an LV end-diastolic dimension >60 mm. What this minority of athletes with pronounced LVH have in common is an enlarged BSA (approximate mean BSA 2.1 m²). However, despite recognising that the largest maximal wall thicknesses are observed in those with the largest BSA’s, there are limited data examining the impact of extreme body anthropometry (BSA>2.3 m²) upon cardiac morphology in professional athletes. This is important as it is widely recognised that LV dimensions are influenced by body anthropometry. The first to recognise the importance of BSA when undertaking preparticipation screening, suggesting that the probability of false-positive HCM identification would be exacerbated in athletes with a BSA>2 m². Basketball, handball and volleyball are three such sports whereby some male athletes may exceed the stereotypical anthropometry for an athlete, with heights and body mass reaching 220 cm and 150 kg, respectively. Magalski et al’s electrocardiographic examination of 1959 American Football players was one of the largest studies to undertake preparticipation screening in athletes with large anthropometry (mean BSA: 2.4±0.3 m²). Regrettably, only 205 American Football players received an echocardiogram following a referral due to an abnormal ECG, family history or clinical examination, with the BSA of this cohort unreported.

The aim of the present study was to investigate the cardiac structure and function in professional male athletes with extreme anthropometry (≥2.3 m²), to confirm if the established upper limits of physiological cardiac adaptation to intensive and sustained physical activity are applicable for this unique population.

METHODS

Ethical approval was obtained from the Shafallah Medical Genetics Centre ethics committee, with all athletes completing informed consent.
Participants
856 asymptomatic professional male athletes (age 25±8 years), exercising ≥6 h/week in 15 high-intensity intermittent sporting disciplines (eg, Soccer, n=586; basketball, n=75; volleyball, n=41 and handball, n=35) presented at our institution for pre-participation cardiac screening (table 1). Athletes were categorised into three distinct groups according to their body surface area

Resting 12-lead ECG
A standard 12-lead ECG was obtained using a GE Mac 5500 (New York, USA) after a period of 5 min rest in the supine position. All ECG’s were reported independently by two experienced investigators (OS and MW) using the recent 2010 ESC recommendations for interpretation of 12-lead electrocardiogram in the athlete, with third opinions sought from two international cardiologists (SS and FC) for difficult cases.

Echocardiography
A two-dimensional (2D), M-mode and Doppler and tissue Doppler echocardiographic examination was performed in the left lateral decubitus position by a consultant cardiologist using a commercially available ultrasound system (Philips, USA). Images of the heart were obtained in the standard parasternal long-axis and short-axis and apical four-chamber planes, as previously described, in order to identify subtle focal regions of SCD. This was important as the population truly reflected those individuals who had not been excluded from competitive sport on the suspicion of harboured an inherited cardiac pathology. As previously described, the term ‘West-Asian’ denotes individuals of Gulf or Middle-Eastern descent, and ‘Black African’ denotes individuals of African descent. West-Asian athletes were recruited from seven Gulf States (Qatar, Bahrain, Oman, UAE, Kuwait, Yemen and Saudi Arabia) and six Middle-Eastern countries (Egypt, Jordan, Palestine, Iraq and Lebanon). Black athletes from seven African countries (Sudan, Somalia, Ghana, Nigeria, Ivory Coast Senegal, Cameroon and Ethiopia) were also recruited alongside Caucasian athletes from the USA, Canada, Australia, Russia, Bosnia and Croatia.

Table 1 Demographic data of all athletes categorised by body surface area (BSA)

Criteria for consideration of the diagnosis of pathological LVH in athletes
On the basis of previous publications and our own experience of an athlete’s heart, athletes with an LV wall thickness >12 mm were considered to have LVH.7 21 22 Athletes with LVH and a relatively non-dilated LV in terms of athletic training (<56 mm)23 in association with any one of the following were considered to have findings that could be consistent with pathological LVH rather than physiological hypertrophy: (1) impaired diastolic function;24 (2) enlarged left atrial diameter >45 mm in athletes
<18-year-old and up to 50 mm in older athletes; (5) LV outflow obstruction; (4) (left bundle branch block and (5) ST-segment depression or deep T wave inversions (<0.2 mV) in ≥2 contiguous anterior, inferior or lateral leads (but not aVR, and III) on the ECG. Athletes demonstrating echocardiographic and/or ECG abnormalities considered to represent pathological LVH were investigated further with 48 h ECG, cardiopulmonary exercise test and cardiac MRI to evaluate the broader phenotype of common cardiomyopathic processes such as HCM and arrhythmogenic right ventricular cardiomyopathy, in addition to assessing the risk of SCD.

Statistical analysis
All data were presented as mean±SD and (range), and analysed using SPSS (Statistical Package for Social Sciences 17, Illinois, USA). Data were analysed using a two-way-between-subjects analysis of variance, with pair-wise comparisons, used to identify any significant differences in athlete anthropometrics and echocardiographic characteristics between the three BSA groups, together with athlete anthropometrics and echocardiographic characteristics between Black African, West-Asian and Caucasian ethnicity in the >2.3 m² BSA group. Relationships between data indices of echocardiographic measures of cardiac structure and function against the athletes BSA together with their resting systolic blood pressure were examined via Pearson’s product–moment correlation analysis. A p value <0.05 was considered significant.

RESULTS
None of the athletes experienced angina, breathlessness disproportionate to the amount of exercise performed, or exertional syncope. The diagnosis of HCM was excluded by echocardiography in 819 (98%) on the basis of an LV wall thickness <12 mm, absence of systolic anterior motion of the anterior mitral valve leaflet causing LV outflow obstruction and normal diastolic function.

Athletes with an LV wall thickness >12 mm (LVH)
Of the 836 athletes, 17 (2%) showed a maximal LV wall thickness exceeding 12 mm and were considered to have LVH. Twelve (12%) of these athletes demonstrating LVH came from the >2.3 m² BSA cohort, compared with just 3 (1.2%) and 2 (0.4%) from Groups 2 and 3, respectively. All 17 athletes with LVH had an appropriate (≥45 mm) LV chamber dimension (mean 56±4 mm, (range 49–65)), normal systolic and diastolic function, an enlarged left atrial diameter and no systolic anterior motion of the anterior mitral valve leaflet or LV outflow obstruction.

Athletes with an LVH >12 mm and an abnormal ECG
Of the 17 athletes with LVH, only 2 (0.2%) demonstrated a wall thickness that exceeded 15 mm (table 2; athletes 1 and 2). However, both athletes also demonstrated ECG’s highly suspicious of HCM (figure 1A,B). Following extensive further investigation, both athletes were diagnosed with a cardiomyopathy and were eventually disqualified from competitive sport. Both athletes were removed from any further group analysis.

Athletes with other cardiac abnormalities on ECG and echocardiography
One further athlete (table 2; athlete 3) was diagnosed with a mild variant of apical HCM, due to disproportionately thickened apical segments of the basal and septal walls upon cardiac MRI, following an abnormal ECG (figure 1C) but normal echocardiogram (maximal basal LV wall thickness of 8 mm). This athlete was removed from any further group analysis. Four asymptomatic athletes without cardiac murmur were found to have trivial valve regurgitation (one mitral, one aortic and two tricuspid), not requiring further investigation after echocardiography. Finally, one asymptomatic player had an aortic root

Table 2 Follow-up results of three athletes presenting with an abnormal ECG on initial screening

<table>
<thead>
<tr>
<th>Athlete</th>
<th>Symptoms</th>
<th>FH of SCD (&lt;35 year)</th>
<th>ECG abnormality</th>
<th>Echocardiogram</th>
<th>Exercise Stress Test and 24 h Holter ECG</th>
<th>CMR</th>
<th>Screened first-degree relatives</th>
<th>Diagnosis</th>
</tr>
</thead>
<tbody>
<tr>
<td>(1) 19-year-old West-Asian footballer (1.99 m² BSA)</td>
<td>No</td>
<td>No</td>
<td>RAE, profound voltage (77 mm), Q waves in II, III, aVF, T wave inversion in I, II, III, aVL, aVF and ST segment depression in II, III and aVF</td>
<td>Subaortic IVSd bulge of 20 mm, without obstruction of the outflow tract</td>
<td>No arrhythmia during exercise with appropriate BP response. Few monomorphic PVB on Holter monitoring</td>
<td>Asymmetric septal hypertrophy with a maximal septal wall thickness of 20 mm versus lateral wall of 11 mm without obstruction. No LGE, oedema or systolic dysfunction</td>
<td>Father’s ECG and Echo confirmed HCM</td>
<td>Non-obstructive HCM</td>
</tr>
<tr>
<td>(2) 29-year-old Black African-American basketball player (2.35 m² BSA)</td>
<td>No</td>
<td>No</td>
<td>Profound voltage in V3, deep T wave inversion in V6</td>
<td>Normal apart from max wall thickness of 13.6 mm</td>
<td>No arrhythmia during exercise with appropriate BP response</td>
<td>Mild asymmetric hypertrophy of IVSd without obstruction (basal 8 mm, mid 15 mm, apical 9 mm), associated with significant mid-septum transmural fibrosis</td>
<td>Not available</td>
<td>Non-obstructive HCM</td>
</tr>
<tr>
<td>(3) 27-year-old West-Asian Futsal player (1.75 m² BSA)</td>
<td>No</td>
<td>No</td>
<td>Profound voltage in V4 (59mm), T wave inversion in II, III, aVF, V2–V6 and ST segment depression in V4–V5</td>
<td>Normal (max wall thickness 8.2 mm)</td>
<td>No arrhythmia during exercise with appropriate BP response</td>
<td>Apical segments are disproportionately thickened, increased basal and septal wall thickness. No LGE, oedema or systolic dysfunction</td>
<td>Not available</td>
<td>Mild variant of apical HCM</td>
</tr>
</tbody>
</table>

BP: blood pressure; BSA: body surface area; CMR: cardiac magnetic resonance imaging; FH: family history; HCM: hypertrophic cardiomyopathy; IVSd, intraventricular septum in diastole; LGE, late gadolinium enhancement; PVB, premature ventricular beats; RAE, right atrial enlargement; SCD, sudden cardiac death.
dimension at the upper limit of normal but given the borderline measure and normality of other parameters, was provided medical clearance and requested to attend yearly echocardiographic examination. These last five athletes remained in the group analysis.

Impact of BSA upon cardiac structure and function
A significant and progressive increase in aortic, left and right atrial, as well as left and right ventricular dimensions and volumes was observed as BSA increased (table 3; p<0.05). No athlete with a normal ECG demonstrated a maximal wall thickness >13 mm and an LVIDd>65 mm. A total of 36 athletes (4% overall) had an LVIDd>60 mm; of which 19 (19%) came from Group 1 compared to 13 (5%) and 4 (0.8%) from Groups 2 and 3, respectively. A significantly higher peak early diastolic mitral flow velocity was observed in Group 3 compared to the two other BSA groups (p<0.05). Furthermore, a peak early mitral annular velocity of the septal wall was significantly lower in Group 1 athletes than the other two BSA groups (p<0.05). No other cardiac functional measures were significantly different between BSA groups. Finally, a significant linear relationship (r=0.3, p<0.0001) was observed between the athletes BSA and their systolic blood pressure (figure 2), and between systolic blood pressure and IVSd (r=0.49), LVIDd (r=0.54), posterior wall thickness in diastole (PWTd) (r=0.51) and LVM (r=0.69, p<0.0001).

Impact of ethnicity upon cardiac structure and function in athletes with a BSA>2.3 m²
Of the 99 remaining athletes in the BSA>2.3 m² cohort (31 West-Asian, 45 Black African and 23 Caucasian athletes), there were no significant differences in BSA between the three ethnicities. Black African athletes had a significantly thicker IVSd and PWTd than both West-Asian and Caucasian athletes (p<0.05). Black African athletes also had significantly larger LA volumes, RA areas and LV masses than West-Asian athletes (p<0.05). West-Asian athletes had a significantly larger LVIDd than Caucasian athletes (p<0.05). There were no significant differences between ethnicities in any diastolic or systolic parameter (table 4).

DISCUSSION
The main finding of the current study was that the upper normal limits for maximal LV dimensions in professional male athletes with extreme anthropometric characteristics (BSA>2.3 m²) remains 14 mm for maximal wall thickness and 65 mm for LV internal diameter during diastole. Furthermore,
for those athletes with the largest BSA's, Black African ethnicity was associated with larger cardiac dimensions than either West-Asian or Caucasian ethnicity. Finally, in a sample of 836 athletes, 3 were diagnosed with a cardiomyopathy; 0.4% prevalence rate that in all cases, the ECG was vital for the initial identification and eventual diagnosis of the disease.

From four large-scale echocardiographic studies examining a total of 5053 elite athletes, only 134 athletes (2.7%) demonstrated a maximal wall thickness greater than 12 mm, with only a further 27 athletes (0.5%) presenting LVH >15 mm. However, out of these 5053 athletes the largest end of range BSA's in all four studies varied from 2.26 to 2.29 m², with only one athlete from...
Whyte et al.\textsuperscript{4} study presenting a BSA of 2.52 m\textsuperscript{2}. This study presents 100 professional male athletes with a BSA >2.3 m\textsuperscript{2} (mean ± 0.1 m\textsuperscript{2}). Only one athlete from this cohort demonstrated LVH>13 mm, but he also exhibited a particular abnormal ECG suggestive of an inherited cardiac disease, and was eventually diagnosed with HCM via cardiac magnetic resonance. Of the remaining 99 asymptomatic athletes, 12 (12%) had a wall thickness greater than 12 mm and 18 (18%) had an LVIDd greater than 60 mm. However, in the presence of a normal ECG, absence of systolic anterior motion of the anterior mitral valve leaflet causing LV outflow obstruction, an appropriately diluted LV, and normal diastolic function, a diagnosis of HCM was excluded in all athletes. This study suggests that irrespective of an athlete’s enlarged BSA, the upper limit of physiological maximal wall thickness remains in the 13–16 mm range. Indeed, our data support the more conservative approach limits suggested by Whyte et al.\textsuperscript{4} such that regardless of extreme body anthropometry, the physiological upper normal limits of LV wall thickness and LVIDd are 14 and 65 mm, respectively.

Despite being referred for echocardiography due to a clinical suspicion of possible cardiac disease based on an abnormal ECG, family history or clinical evaluation, two studies are worthy of mention that support our data in ‘big’ athletes. Abernethy et al.\textsuperscript{12} investigated 156 asymptomatic American Football (NFL) players and reported mean maximal wall thickness and LVIDd to be 11.2±0.2 and 55±0.5 mm, respectively. While Magalski et al.\textsuperscript{11} observed that from 203 referred NFL athletes, 197 (97%) demonstrated a maximal IVSd from 7 to 12 mm, with six athletes (3%) presenting an IVSd from 13 to 14 mm. However, all 203 were considered not to have a cardiomyopic process due to no other echocardiographic abnormalities and normal Doppler inflow velocities.

It is well recognised that little account of body size is taken in the determination of ‘apparently normal’ cardiac dimensions in adult athletes, even though allometric scaling in the paediatric population is routine practice.\textsuperscript{26} Despite the strong relationships between BSA and multiple LV measures in the current study, absolute upper normal limits are still clinically relevant. It may be that scaling, via an appropriate method and scaling variable, maybe be of more clinical value in those athletes with smaller BSA’s.

Recent data suggest that an athlete’s ethnic origin may have a significant impact on their cardiovascular response to exercise\textsuperscript{13} 35 37 Despite no significant difference in BSA between ethnicities in the >2.3 m\textsuperscript{2} cohort of athletes, Black African athletes presented significantly greater wall thicknesses, and resultant LV masses, than West-Asian and Caucasian athletes. It should be noted that regardless of ethnicity and the extreme BSA of this cohort, the established upper limits of cardiac structure and function are applicable to all three ethnicities. Previous data from our group have demonstrated that a minority

Table 4 Impact of ethnicity upon cardiac structure and function in athletes with a BSA>2.3m\textsuperscript{2} (mean±SD; range)

<table>
<thead>
<tr>
<th></th>
<th>West-Asian</th>
<th>Black African</th>
<th>Caucasian</th>
</tr>
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<tbody>
<tr>
<td>Height (cm)</td>
<td>189±8</td>
<td>201±7</td>
<td>199±7</td>
</tr>
<tr>
<td></td>
<td>(170–206)</td>
<td>(182–217)</td>
<td>(180–210)</td>
</tr>
<tr>
<td>Body mass (kg)</td>
<td>109±14</td>
<td>105±12</td>
<td>101±6</td>
</tr>
<tr>
<td></td>
<td>(91–150)</td>
<td>(85–156)</td>
<td>(87–117)</td>
</tr>
<tr>
<td>Body surface area (m\textsuperscript{2})</td>
<td>2.4±0.1</td>
<td>2.4±0.1</td>
<td>2.4±0.1</td>
</tr>
<tr>
<td></td>
<td>(2.3–2.8)</td>
<td>(2.3–3.0)</td>
<td>(2.3–2.6)</td>
</tr>
<tr>
<td>Aortic diameter (mm)</td>
<td>30±2</td>
<td>30±2</td>
<td>30±2</td>
</tr>
<tr>
<td></td>
<td>(25–33)</td>
<td>(26–35)</td>
<td>(26–34)</td>
</tr>
<tr>
<td>LA (mm)</td>
<td>37±4</td>
<td>37±4</td>
<td>37±4</td>
</tr>
<tr>
<td></td>
<td>(30–45)</td>
<td>(26–45)</td>
<td>(29–48)</td>
</tr>
<tr>
<td>LA area (mm\textsuperscript{2})</td>
<td>22±3</td>
<td>23±3</td>
<td>22±4</td>
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<tr>
<td></td>
<td>(14–30)</td>
<td>(16–29)</td>
<td>(14–33)</td>
</tr>
<tr>
<td>LA volume (ml)</td>
<td>61±15</td>
<td>71±16**</td>
<td>60±18</td>
</tr>
<tr>
<td></td>
<td>(23–91)</td>
<td>(38–103)</td>
<td>(31–98)</td>
</tr>
<tr>
<td>Right atrial area (mm\textsuperscript{2})</td>
<td>19±4</td>
<td>21±3**</td>
<td>19±3</td>
</tr>
<tr>
<td></td>
<td>(12–28)</td>
<td>(12–28)</td>
<td>(13–24)</td>
</tr>
<tr>
<td>Interventricular septum thickness in diastole (mm)</td>
<td>10±1</td>
<td>11±1*</td>
<td>10±1</td>
</tr>
<tr>
<td></td>
<td>(7–12)</td>
<td>(6–13)</td>
<td>(7–13)</td>
</tr>
<tr>
<td>LVID in diastole (mm)</td>
<td>60±1***</td>
<td>57±1</td>
<td>58±1</td>
</tr>
<tr>
<td></td>
<td>(48–80)</td>
<td>(50–63)</td>
<td>(54–63)</td>
</tr>
<tr>
<td>Posterior wall thickness in diastole (mm)</td>
<td>9±0.2</td>
<td>10±0.2*</td>
<td>9±0.2</td>
</tr>
<tr>
<td></td>
<td>(7–12)</td>
<td>(7–13)</td>
<td>(8–10)</td>
</tr>
<tr>
<td>LVID in systole (mm)</td>
<td>39±5</td>
<td>40±5</td>
<td>41±3</td>
</tr>
<tr>
<td></td>
<td>(28–46)</td>
<td>(26–50)</td>
<td>(36–46)</td>
</tr>
<tr>
<td>LV end diastolic volume (ml)</td>
<td>147±32</td>
<td>157±24</td>
<td>161±36</td>
</tr>
<tr>
<td></td>
<td>(88–236)</td>
<td>(109–209)</td>
<td>(120–271)</td>
</tr>
<tr>
<td>LV end systolic volume (ml)</td>
<td>48±12</td>
<td>51±10</td>
<td>54±16</td>
</tr>
<tr>
<td></td>
<td>(22–76)</td>
<td>(34–75)</td>
<td>(30–111)</td>
</tr>
<tr>
<td>LV mass (g)</td>
<td>208±33</td>
<td>236±43**</td>
<td>217±27</td>
</tr>
<tr>
<td></td>
<td>(140–276)</td>
<td>(146–348)</td>
<td>(164–266)</td>
</tr>
</tbody>
</table>

*Significant difference between Black athletes versus West-Asian and Caucasian athletes (p<0.05).
**Significant difference between Black athletes versus West-Asian athletes (p<0.05).
***Significant difference between West-Asian athletes versus Caucasian athletes (p<0.05).

LA, left atrial diameter; LVIDd, left ventricular internal diameter.
(3%) of Black athletes (mean BSA 2.1 m²; 9 from 300) may present physiological LVH [34]. The present data set of 219 Black African athletes from a broad range of BSA's did not find an athlete with a normal ECG presenting a maximal wall thickness greater than 15 mm. A limitation of the present study is that a data set of 46 Black African with a BSA>2.3 m² maybe too small to ascertain if this ethnicity with extreme anthropometry requires the upper limits of physiological LVH to be raised to 15 mm; in spite of demonstrating significantly greater wall thicknesses and masses that their West-Asian and Caucasian counterparts. Interestingly, Basavarajaiah et al [35] study demonstrated that basal and exercise-related BP responses to physical activity are greater wall thicknesses and masses that their West-Asian and Caucasian counterparts. Interestingly, Basavarajaiah et al study demonstrated that basal and exercise-related BP responses to physical activity are.

Regardless of the increased LVH in black athletes. Yet to date, no data have been published to confirm this postulation. Second, while professional basketball, handball and volleyball players were recruited, substantiation of the upper limits of physiological LVH is required in the small minority of athletes with extreme BSA's who compete in sports that induce the greatest amounts of cardiac remodelling; namely rowing, cycling, cross-country skiing, biathlon. Nevertheless, this will once again prove problematic for the Black African ethnicity whose participation in these endurance sports is limited.

Regardless of these limitations, our data support the clinical utility of ECG in the initial identification of athletes suspected of harbouring an inherited cardiac disease. The present study diagnosed one athlete with a mild variant of apical HCM via cardiac magnetic resonance, following an abnormal ECG suggestive of an inherited cardiac disease. This was in spite of a normal echo-cardiogram (maximal LV wall thickness of 8 mm). In conclusion, marked repolarisation changes, ST depression, pathological Q waves and multiple ventricular ectopic's are a great concern, even when cardiac dimensions are within accepted limits.

In conclusion, irrespective of an athlete’s extreme anthropometrical dimensions and ethnicity, the physiological upper normal limits of LV wall thickness and LV internal diameter during diastole due to intensive and sustained physical activity are 14 and 65 mm, respectively. Moreover, even when matched for extreme BSA (>2.3 m²), Black African athletes present significantly greater wall thicknesses and resultant LV masses than Arabic and Caucasian athletes.

What this study adds

- Regardless of extreme body surface area (BSA), no asymptomatic athlete with a negative family history of sudden cardiac death and a normal ECG presented a maximal wall thickness >14 mm.
- Established upper limits for physiological cardiac hypertrophy of 14 mm for maximal wall thickness and 65 mm for left ventricular internal diameter during diastole are appropriate, irrespective of an athlete’s BSA.
- We identified one athlete with a cardiomyopathy with normal wall thicknesses on echocardiography but abnormal cardiac magnetic resonance. The ECG remains key and should guide follow-up management. Marked repolarisation changes, ST depression, pathological Q waves and multiple ventricular ectopic’s are a great concern, even when cardiac dimensions are within accepted limits on echocardiography.

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