Run for your life ... at a comfortable speed and not too far

James H O’Keefe,1,2 Carl J Lavie3,4

During the Greco-Persian War in 490 BCE, Philipppides, a 40-year-old herald messenger (professional running-courier) ran the 26 miles from a battlefield near Marathon, Greece, into Athens carrying momentous news of Greek victory. Upon arriving at the Acropolis, he proclaimed: ‘Joy, we have won!’ and then immediately collapsed and died.1 Fast-forward about 2500 years to an era when the baby-boomer’s came of age and long-distance running boomed. The prevailing logic held that aerobic exercise is clearly good for one’s health and that, if some is good, more must be better. In 1975, Dr Thomas Bassler, a physician/runner, boldly proclaimed that, if you could run a marathon, you were immune to death from coronary heart disease (CHD).2 This urban myth has long since been disproven; indeed an emerging body of evidence suggests the opposite: extreme endurance exercise may exact a toll on cardiovascular (CV) health.

'SHOW ME THE BODIES'

After our recent articles on this topic,1 3–5 Ambry Burfoot, winner of the 1968 Boston Marathon and Editor-at-Large for Runner’s World Magazine, challenged our assertions about the dangers of extreme endurance efforts by demanding, ‘Show me the bodies’. Ambry has a good point: the risk of dropping dead in a marathon is remote, about 0.5 to 1 in 100,000 participants.6 But the occasional marathoner or triathlete who dies while strenuously exercising is the ‘canary in the coal mine’. Chronic extreme exercise appears to cause excessive ‘wear-and-tear’ on the heart, inducing adverse structural and electrical remodelling, which offsets some of the CV benefits and longevity improvements conferred by moderate physical activity. Thus, even though chronic extreme exercise may not kill you, it may erase many of the health advantages of regular moderate exercise.

Indeed, regular vigorous exercise is probably the single best step a person can take to ensure robust CV health. In a study of 416 000 adults followed for a mean of 8 years, 40–50 min per day of vigorous exercise reduced risk of death by about 40% (figure 1).7 In that study, at about 45 min, a point of diminishing returns is reached whereby longer exercise efforts do not appear to translate into lower death risk. Light to moderate physical activity reduced death rates too, albeit not as strongly, but in this case more physical activity appeared to be better, with no plateau out to 110 min daily. Indeed, if we had a pill that confers all the benefits of exercise, many physicians might be looking for work. Approximately 30–45 min of daily vigorous exercise significantly reduces risks for many maladies including early death, Alzheimer’s disease, CHD, diabetes, osteoporosis and depression.4 5 Yet, as can be expected with any potent drug, an insufficient dose will not confer the optimal benefits, while an excessive dose can cause harm, and even death in extreme overdoses.

The ‘survival of the fittest’ concept does not fully apply to the modern world, where it appears that even the moderately fit have an excellent CV prognosis and superb longevity. Studies of CV fitness, as measured by peak performance on a treadmill, show a curvilinear relationship whereby improvements from unfit to moderately fit confer dramatic reductions in morbidity and mortality (figure 2).8 However, fitness levels above 12 metabolic equivalents do not seem to translate into additional gains in CV health and longevity. Thus, if one is training to be able to run at speeds above 7.5 miles per hour, this is being done for some reason other than further improvements in life expectancy.

CV DAMAGE FROM EXCESSIVE EXERCISE

High-intensity exercise sessions lasting beyond 1–2 h cause acute volume overload of the atria and right ventricle (RV), which can bring about overstretching and micro-tears in the myocardium, as evidenced by a transient rise in cardiac biomarkers, including troponin and B-natriuretic peptide and a fall in the RV ejection fraction.9 Although within 1 week, these transitory abnormalities usually return to baseline,9 after years to decades of excessive exercise and repetitive injury, this pattern can lead to patchy myocardial fibrosis, particularly in the pliable walls of the heart such as the atria and RV, creating a substrate for atrial and potentially malignant ventricular arrhythmias.1 3–5 In addition, long-term excessive exercise may accelerate aging in the heart, as evidenced by increased coronary artery calcification, diastolic ventricular dysfunction, and large-artery wall stiffening.1 3–5

At rest, the heart pumps about 5 litres/min; with strenuous aerobic exercise, the cardiac output can rise 5–7-fold, pumping up to 25–35 litres/min. This massive increase in cardiac work is what the heart is designed to do for short bursts, or even for up to as long as 30 or 50 min
continuously. However, with protracted efforts, these high volumes can overstretch the chambers, eventually disrupting cardiac muscle fibres and causing micro-tears in the myocardium. The presence of sustained exercise-induced elevations in catecholamines and pro-oxidant free radicals worsen the situation by adding inflammation to the injury, leading eventually to scarring and stiffening of the CV structures.

A trial randomised 60 male patients with CHD to vigorous exercise sessions of either 30 or 60 min. The 30 min exercise workouts improved arterial elasticity and produced minimal oxidant stress. In contrast, the 60 min sessions increased oxidant stress and worsened vascular stiffness as measured by pulse wave velocity, particularly in those over the age of 50. MRI scans of runners who have been participating in marathons for decades show a threefold increased incidence of scattered fibrosis and scarring in the walls of the atria, interventricular septum and RV (figure 3).

Cardiologists from Minnesota evaluated a group of runners who had completed at least 25 marathons over 25 years and found a 60% increase in coronary plaque burden compared with sedentary age-matched controls. These findings were replicated by a group from Germany, who showed increased coronary plaque in 108 chronic marathoners compared with sedentary controls. This scarring can set the stage for dangerous heart rhythms, such as atrial fibrillation, which is increased approximately fivefold in veteran endurance athletes. Ventricular tachycardia and sudden cardiac arrest can also be seen in endurance athletes even in the absence of CHD and hypertrophic cardiomyopathy.

An enlightening study by Benito et al reinforced the concept of cardiac damage from chronic excessive exercise. Mice after being forced to run to exhaustion every day for 4 months showed the same cardiac enlargement, scarring and predisposition to dangerous ventricular dysrhythmias that have been documented in some veteran extreme endurance athletes. Encouragingly, when the mice were withdrawn from the ‘Iron-Mouse’ training regimen and allowed to resume normal mouse physical activity levels, their cardiac abnormalities showed marked improvements, even showing regression of myocardial fibrosis and resolution of the tendency toward serious ventricular dysrhythmias.

**PHIDIPPIDES CARDIOMYOPATHY**

*Born to Run* is a non-fiction bestseller book published in 2009 that glamorises ultra-endurance running. The story’s hero is Micah True, an American who dropped out of modern civilisation to live and run with the Tarahumara Indians in Mexico. Nicknamed Caballo Blanco, or white horse, for his legendary running endurance, he routinely ran daily distances of 25–100 miles. This March on a 12-mile training run in New Mexico, Micah True dropped dead at age 58. On autopsy, his heart was enlarged and thickened with ‘focal areas of interstitial chronic inflammatory infiltrate’ in the myocardium; the coronary arteries were ‘focally thickened with mild coronary arteriosclerosis’. Chief Medical Investigator Ross Zumwalt, MD summarised the findings as, ‘Unclassified cardiomyopathy, which resulted in a cardiac dysrhythmia during exertion’. When considered in the context of True’s decades-long lifestyle of daily ultra-marathon running, we suspect that the autopsy findings were an example of ‘Phidippides cardiomyopathy’—the constellation of cardiac pathology that has been in observed in the hearts of some veteran extreme endurance athletes.

![Figure 2](https://example.com/image1)

**Figure 2** Death rates as a function of cardiovascular fitness as measured by metabolic equivalents achieved on maximal exercise treadmill testing. CVD, cardiovascular disease.

![Figure 3](https://example.com/image2)

**Figure 3** MRI scans showing scattered scarring (red arrows) in the heart, especially in the interventricular septum.
MORONIC EXERCISE: THE SWEET-SPOT FOR LONGEVITY

Two very recent studies presented in abstract form at major national meetings may revolutionise our thinking about running and its health effects. One is a prospective observational study that followed 52 600 people for up to three decades. The 14 000 runners in that study had a 19% lower risk of death compared with the 42 000 non-runners. Yet, when they sub-grouped the runners by weekly mileage, those who ran over 20 or 25 miles per week seemed to lose their survival advantage over the non-runners (figure 4). On the other hand, those who ran between 5 and 20 miles total per week enjoyed a 25% decrease in risk of death during follow-up. The same pattern emerged for speed of running: the bottom of the U curve—moderate exercise—is the ‘sweet spot’ for which most should try to aim. Sitting is the new smoking; a sedentary lifestyle will cause disability and disease, and will shorten life expectancy. We are not so much born to run as born to walk. Ethnographic research indicates that, in the environment of human evolution, our ancient ancestors walked 4–10 miles a day.1 Walking is superior to running for mechanical efficiency and musculoskeletal durability. Indeed, we advise our patients that they can walk or garden hours a day without concern about CV overuse injury.

So while it is true that exercise confers powerful health benefits, the common belief that more is better is clearly not true. The unique and potent benefits of exercise are best bestowed by moderate exercise and physical activity. The exercise patterns for maximising CV fitness/peak aerobic capacity are very different from those that best confer CV health, durability and overall longevity. So, if one’s goal in life is to compete in the marathon or triathlon of the Rio Olympics in 2016, this will certainly require high-intensity exercise for hours a day. But, for those whose goal is to be alive and well while watching the 2052 Olympics from the stands, exercise and physical activity at lower intensities and durations would be more ideal.

CONCLUSION

The take home message for most is to limit one’s vigorous exercise to 30–50 min/day. If one really wants to do a marathon or full-distance triathlon etc, it may be best to do just one or a few and then proceed to safer and healthier exercise patterns. On the other hand, light or moderate intensity exercise does not present the dose-dependent risks associated with excessive endurance exercise. A routine of moderate physical activity will add life to your years, as well as years to your life. In contrast, running too fast, too far, and for too many years may speed one’s progress towards the finish line of life.

Contributors JHO and CJL contributed.

Competing interests None.

Provenance and peer review Commissioned; internally peer reviewed.

Author note A video presentation on this topic is available on the internet: YouTube, TEDx Talk, James O’Keefe, Run for your life… at a comfortable pace and not too far.

Heart 0:0 1–4. doi:10.1136/heartjnl-2012-302886

REFERENCES

9. Ector J, Ganneau J, van der Merwe N, et al. Reduced right ventricular ejection fraction in

THE U-CURVE

Hippocrates, the father of medicine and a contemporary of Phidippides in ancient Greece, taught, ‘The right amount of nourishment and exercise, not too much, not too little, is the safest way to health’.1 If you listen to your body, this is just common sense. Yet, nothing we have published previously has stirred so much controversy, especially among the general public. Increasingly our culture is one of extremes: during the past 30 years, obesity has tripled in the USA and has increased in much of the Western World, while during the same time the number of people completing a marathon has risen 20-fold. On one side of the U-curve, the couch loungers/channel surfers embrace this message as justification for continuing their sedentary lifestyle. And, on the far end of the U-curve, the extreme exercise aficionados want to ignore the message and instead kill the messenger. As with many things in life, the safe and comfortable zone at the bottom of the U curve—moderate exercise—is the ‘sweet spot’ for which most should try to aim.

Figure 4 All-cause mortality by running distance per week.16


Run for your life … at a comfortable speed and not too far

James H O'Keefe and Carl J Lavie

*Heart* published online November 29, 2012
doi: 10.1136/heartjnl-2012-302886

Updated information and services can be found at:
http://heart.bmj.com/content/early/2012/11/21/heartjnl-2012-302886.full.html

**These include:**

**Data Supplement**

"Press release"
http://heart.bmj.com/content/suppl/2012/12/11/heartjnl-2012-302886.DC1.html

"HTML Page - index.html"
http://heart.bmj.com/content/suppl/2013/01/09/heartjnl-2012-302886.DC2.html

**References**

This article cites 14 articles, 5 of which can be accessed free at:
http://heart.bmj.com/content/early/2012/11/21/heartjnl-2012-302886.full.html#ref-list-1

**P<box>**

Published online November 29, 2012 in advance of the print journal.

**Email alerting service**

Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

**Topic Collections**

Articles on similar topics can be found in the following collections

- Press releases (17 articles)
- Epidemiology (2566 articles)
- Drugs: cardiovascular system (6394 articles)

Advance online articles have been peer reviewed, accepted for publication, edited and typeset, but have not yet appeared in the paper journal. Advance online articles are citable and establish publication priority; they are indexed by PubMed from initial publication. Citations to Advance online articles must include the digital object identifier (DOIs) and date of initial publication.

To request permissions go to:
http://group.bmj.com/group/rights-licensing/permissions

To order reprints go to:
http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to:
http://group.bmj.com/subscribe/
Advance online articles have been peer reviewed, accepted for publication, edited and typeset, but have not yet appeared in the paper journal. Advance online articles are citable and establish publication priority; they are indexed by PubMed from initial publication. Citations to Advance online articles must include the digital object identifier (DOIs) and date of initial publication.

To request permissions go to:
http://group.bmj.com/group/rights-licensing/permissions

To order reprints go to:
http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to:
http://group.bmj.com/subscribe/
Potential Adverse Cardiovascular Effects From Excessive Endurance Exercise

James H. O’Keefe, MD; Harshal R. Patil, MD; Carl J. Lavie, MD; Anthony Magalski, MD; Robert A. Vogel, MD; and Peter A. McCullough, MD, MPH

Abstract

A routine of regular exercise is highly effective for prevention and treatment of many common chronic diseases and improves cardiovascular (CV) health and longevity. However, long-term excessive endurance exercise may induce pathologic structural remodeling of the heart and large arteries. Emerging data suggest that chronic training for and competing in extreme endurance events such as marathons, ultramarathons, ironman distance triathlons, and very long distance bicycle races, can cause transient acute volume overload of the atria and right ventricle, with transient reductions in right ventricular ejection fraction and elevations of cardiac biomarkers, all of which return to normal within 1 week. Over months to years of repetitive injury, this process, in some individuals, may lead to patchy myocardial fibrosis, particularly in the atria, interventricular septum, and right ventricle, creating a substrate for atrial and ventricular arrhythmias. Additionally, long-term excessive sustained exercise may be associated with coronary artery calcification, diastolic dysfunction, and large-artery wall stiffening. However, this concept is still hypothetical and there is some inconsistency in the reported findings. Furthermore, lifelong vigorous exercisers generally have low mortality rates and excellent functional capacity. Notwithstanding, the hypothesis that long-term excessive endurance exercise may induce adverse CV remodeling warrants further investigation to identify at-risk individuals and formulate physical fitness regimens for conferring optimal CV health and longevity.

© 2012 Mayo Foundation for Medical Education and Research  Mayoclinicproceedings.org

From Mid America Heart Institute of Saint Luke’s Hospital of Kansas City, MO (J.H.O., H.R.P., A.M.); John Ochsner Heart and Vascular Institute, Ochsner Clinical School–The University of Queensland School of Medicine, New Orleans, LA, and Pennington Biomedical Research Center, Louisiana State University System, Baton Rouge (C.J.L.); University of Maryland, Baltimore (R.A.V.); and St. John Providence Health System Providence Park Heart Institute, Novi, MI (P.A.M.).
SUDDEN CARDIAC DEATH AND ENDURANCE ET

Over the past 35 years, the number of Americans participating in a marathon annually has risen 20-fold; in 2010, an estimated half-million runners completed a marathon in the United States.10 Sudden cardiac death (SCD) among marathoners is very rare, with 1 event per 100,000 participants.6,7,11,12 Although that per-participant risk has not changed over the decades, absolute mortality rates have increased as the number of participants has risen. The final 1 mile of the marathon course represents less than 5% of the total distance of 26.2 miles yet accounts for almost 50% of the SCDs during the race.12,13

The fatality rate for triathlons is approximately twice that of marathons, largely because of increased CV events and drownings during the swim portion of the races.14 The incidence of SCD among collegiate athletes during competition is about 1 per 40,000 participants per year for all athletes.15 It is extremely important to keep in mind that the occurrence of SCD during marathons, triathlons, and collegiate athletic events is rare and should not deter individuals from participating in vigorous ET; the benefits of regular PA to the individual and to society as a whole far outweigh potential risks. At the same time, long-term training for and competing in extreme endurance events may predispose to CV issues that are not seen in more moderate forms of PA.

The causes of SCD during or after extreme exertion in individuals younger than 30 years most commonly include genetic causes such as hypertrophic cardiomyopathy, anomalous coronary arteries, dilated cardiomyopathy, and congenital long QT syndrome. In athletes older than 30 years, CHD and acute myocardial infarction16 and ischemia are the predominant causes of exercise-related SCD.17-23

ANIMAL STUDIES

In an elegant animal model of excessive endurance ET, rats were trained (in part by prodding with electrical shocks to maintain high-intensity effort) to run strenuously and continuously for 60 minutes daily for 16 weeks, and then they were compared with control sedentary rats.8,24 The running rats developed hypertrophy of the left ventricle (LV) and the right ventricle (RV), diastolic dysfunction, and dilation of the left atria and the right atria (RA); they also showed increased collagen deposition and fibrosis in both the atria and ventricles (Figure 2). Ventricular tachycardia was inducible in 42% of the

ARTICLE HIGHLIGHTS

- People who exercise regularly have markedly lower rates of disability and a mean life expectancy that is 7 years longer than that of their physically inactive contemporaries. However, a safe upper-dose limit potentially exists, beyond which the adverse effects of exercise may outweigh its benefits.
- Chronic intense and sustained exercise can cause patchy myocardial fibrosis, particularly in the atria, interventricular septum, and right ventricle, creating a substrate for atrial and ventricular arrhythmias.
- Chronic excessive sustained exercise may also be associated with coronary artery calcification, diastolic dysfunction, and large-artery wall stiffening.
- Veteran endurance athletes in sports such as marathon or ultramarathon running or professional cycling have been noted to have a 5-fold increase in the prevalence of atrial fibrillation.
- Intense endurance exercise efforts often cause elevation in biomarkers of myocardial injury (troponin and B-type natriuretic peptide), which were correlated with transient reductions in right ventricular ejection fraction.

TABLE 1. Potential Benefits of Exercise Training

<table>
<thead>
<tr>
<th>Category</th>
<th>Benefits</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coronary heart disease risk factors</td>
<td>Increases serum high-density lipoprotein cholesterol levels, reduces serum triglyceride and possibly low-density lipoprotein cholesterol levels, reduces indices of obesity, reduces arterial blood pressure, improves insulin sensitivity and glucose levels, improves endothelial function, helps with smoking cessation efforts, reduces psychological stress</td>
</tr>
<tr>
<td>Hematologic</td>
<td>Decreases hematocrit and blood viscosity, expands blood plasma volume, increases red blood cell deformability and tissue-level perfusion, increases circulatory fibrinolytic activity</td>
</tr>
<tr>
<td>Other</td>
<td>Increases coronary flow reserve, increases coronary collateral circulation, increases tolerance of ischemia, increases myocardial capillary density, increases ventricular fibrillation thresholds, reduces atherosclerosis, possibly increases epicardial coronary artery size, reduces major morbidity and mortality</td>
</tr>
</tbody>
</table>

From Mayo Clin Proc.4
running rats vs only 6% of the sedentary rats (P = .05). Importantly, the fibrotic changes caused by 16 weeks of intensive ET had largely regressed to normal by 8 weeks after the daily running regimen ceased.

This animal study found that daily excessive, strenuous, uninterrupted running replicated the adverse cardiac structural remodeling and proarrhythmia substrate noted in observational studies of extreme endurance athletes. These findings support the hypothesis that in some individuals, long-term strenuous daily endurance ET, such as marathon running or professional long-distance cycling, in some individuals may cause cardiac fibrosis (especially in the atria and the RV and interventricular septum), diastolic dysfunction, and increased susceptibility to atrial and ventricular arrhythmias. Many previous animal studies have also found acute, adverse cardiac effects of prolonged (up to 6 hours) endurance exercise, sometimes employing a rat model of cold-water swimming in which the animals were forced to swim to avoid drowning. These studies are of uncertain clinical relevance because of the excessively stressful nature of the imposed exercise.

**ATHLETE’S HEART**

Chronic ET imposes increased hemodynamic demands that alter the loading conditions of the heart, particularly among athletes participating in sports requiring sustained elevations in cardiac work, such as long-distance running, rowing, swimming, and cycling. Highly trained individuals develop cardiac adaptations including enlarged LV and RV volumes, increased LV wall thickness and cardiac mass, and increased left atrial size. In the general population, these structural changes are associated with poor cardiac prognosis. However, these structural alterations, together with a preserved LV ejection fraction (EF), have been considered typical findings of the “athlete’s heart.” Of concern, accumulating information suggests that some of the remodeling that occurs in endurance athletes may not be entirely benign. For example, in elite athletes, cardiac dimensions do not completely regress to normal levels even several years after the athlete has retired from competition and heavy ET.

**BIOMARKER EVIDENCE FOR CARDIAC DAMAGE WITH EXTREME ENDURANCE ET**

Running is a prototypical natural PA and often plays an integral and important role in an active, healthy lifestyle. However, uninterrupted very long-distance running as is generally done while training for and participating in marathons and other extreme endurance events may produce adverse CV effects in susceptible individuals. Serologic markers of cardiac damage, including cardiac troponin, creatine kinase MB, and B-type natriuretic peptide, have been documented to increase in up to 50% of participants during and after marathon running (Figure 3). Additionally, transient renal dysfunction has been observed with extreme endurance ET efforts causing volume depletion and diminished renal filtration, with elevations in serum urea nitrogen, serum creatinine, and cystatin C. Increased levels of cardiac biomarkers including troponin after extreme ET endurance events, such as marathons, may reflect myocardial cell damage at the sites of myocyte slippage of one cell along another due to loss of integrity of desmosomal connections. However, the significance of the elevated cardiac biomarkers after endurance efforts remains uncertain, and it has been argued that these may be entirely benign transient increases resulting from CV adaptations to long-term ET.

**ADVERSE STRUCTURAL REMODELING**

Accumulating evidence suggests that the adverse effects of both short-term intense PA and cumulative endurance exercise are most apparent in the right-sided cardiac chambers. Cardiac output at rest is approximately 5 L/min but typically increases 5-fold to about 25 L/min during vigorous ET. Long-term daily sessions of hours of continuous strenuous PA cause dilation of the RA and RV. During the postexercise period, the cardiac geometric dimensions are restored, but with this recurrent stretch of the chambers and reestablishment of the chamber geometry, some individuals may be prone to the development of chronic structural changes including chronic dilatation of the RV and RA with patchy myocardial scarring in response to the recurrent volume overload.
load and excessive cardiac strain. These abnormalities are often asymptomatic and probably accrue over many years; they might predispose to serious arrhythmias such as atrial fibrillation and/or ventricular arrhythmias (VAs).

A prospective study of 25 runners (13 women and 12 men) found that running a marathon caused acute dilation of the RA and RV, with a sudden decrease in the RVEF. La Gerche et al studied a cohort of 40 highly trained aerobic athletes after competing in endurance events including marathon (mean time to completion, 3 hours), half-ironman triathlon (5.5 hours), full-ironman triathlon (11 hours), and alpine bicycle race (8 hours). They found that these intense endurance exercise efforts caused elevations in biomarkers of myocardial injury (troponin and B-type natriuretic peptide), which were correlated with reductions in RVEF (Figure 4), but not LVEF, on immediate (mean, 45 minutes) post-race echocardiography. The reductions in RVEF and the increases in RV volumes, which returned entirely to baseline within 1 week, were seen most often in races of longer durations (Figure 5). Of this cohort of endurance athletes, 5 (12.5%) had myocardial scarring as detected by local gadolinium enhancement on cardiac magnetic resonance imaging (MRI) (Figure 6). The myocardial scarring and chronic RV remodeling were more common in athletes with the largest cumulative experience in competitive endurance events.

In summary, this study suggests that intense endurance exercise induces acute RV dysfunction while largely sparing the LV. Even when short-term RV recovery appears complete, long-term training for and competing in extreme endurance exercise may lead to myocardial fibrosis and remodeling in a small subgroup.

Ector et al reported that the decrease in RVEF is less significant in athletes with no symptoms of arrhythmia than in endurance athletes who have symptoms of arrhythmia, in whom the RV size increases and the RVEF is significantly lower. Another study of endurance athletes who have symptoms of VAs found that 50% of them had RV structural ab-
normalities by MRI. This RV dysfunction is likely induced by recurrent extreme and sustained high-level PA, with marked elevations in pulmonary artery pressures of up to 80 mm Hg in some athletes, which eventually may cause scattered areas of myocardial injury (as evidenced by the increases in troponin) with subsequent fibrotic scarring, typically in the RV and atria. These observations have led to speculation about the existence of a syndrome of exercise-induced arrhythmogenic RV cardiomyopathy that shares some features with the familial RV disease but is caused by chronic high-level endurance ET rather than a genetic predisposition.

Another study using MRI to assess the effects of long-term very long distance running on myocardial structure comprised 102 ostensibly healthy male runners ranging in age from 50 to 72 years who had completed at least 5 marathons during the previous 3 years, compared with 102 age-matched controls. Approximately 12% of these apparently healthy marathon runners have evidence of patchy myocardial scarring, manifested as late gadolinium enhancement; this was 3-fold more common than in age-matched controls. Of additional concern, the CHD event rate during 2-year follow-up was significantly higher in the marathon runners than in controls (P<0.001). A similar smaller study found pathologic myocardial fibrosis by cardiac MRI in 6 of 12 asymptomatic men (50%) who were lifelong veteran endurance athletes, but no cases in younger endurance athletes and age-matched controls.

Aortic stiffness and arterial pulse wave velocity, which are markers for adverse CV prognosis, may be increased in veteran ultraendurance athletes. A study of 47 individuals who trained extensively for and competed in marathons found that pulse wave velocity and aortic stiffness were significantly higher in the group of marathoners compared with controls. It is possible that the sustained shear stress caused by protracted endurance efforts eventually may induce fibrotic changes and decreases in arterial wall elasticity. Diastolic dysfunction of both the RV and LV has also been observed in individuals doing long-term extreme ET and racing.

CORONARY ARTERY CHANGES

Veteran endurance marathon runners in one study had coronary arteries that, at resting baseline, were similar in size to those of sedentary controls, but the marathoners had greater coronary artery dilating capacity. Mohlenkamp et al studied 108 middle-aged German long-term marathon runners and compared them with matched nonrunner controls. They observed a greater atherosclerotic burden in the marathoners as documented by higher coronary artery calcium (CAC) scores. Additionally, during follow-up the adverse CV event rates in the marathoners were equivalent to those in a population with established CHD. In a similar study, Schwartz et al reported on a US cohort of long-term marathon runners, defined as individuals who completed at least 25 marathons over the previous

![FIGURE 3. High-sensitivity cardiac troponin T (hs-cTnT) concentrations before, immediately after, and 24 and 72 hours after marathon race. From Med Sci Sports Exerc, with permission.](image-url)

![FIGURE 4. Duration-dependent effect of endurance events on right ventricular (RV) ejection fraction. From Eur Heart J, with permission.](image-url)
25 years, and found higher than expected levels of CAC and calcified coronary plaque volume. That study, utilizing computed tomographic coronary angiography, found that the long-term marathoners had significantly more calcified plaque volume than sedentary controls (mean, 274 mm³ vs 169 mm³).

In a case report, Goel et al.54 observed a 49-year-old marathoner who had significant obstructions in all 3 major epicardial coronary arteries without associated risk factors and who generated protracted oxidative stress with prolonged running.

In another study of veteran endurance athletes, mean LV mass, as determined by MRI, was significantly greater in a group of marathon runners than in controls, and the increased LV mass correlated with higher CAC scores. Specifically, those marathoners with an LV mass greater than 150 g had a significantly higher CAC score than those with an LV mass less than 150 g.30 The investigators also found a mismatch between the risk factor profile and the amount CAC, particularly in the marathoners with an LV mass greater than 150 g.

**PATHOPHYSIOLOGY OF LONG-TERM EXTREME ET**

Figure 7 shows the pathophysiology and possible adverse CV consequences (fibrosis, atrial arrhythmias, VAs, and SCD) associated with endurance ET and competition, such as marathon running. Individuals who do long-term ET and race over very long distances induce sustained (often for 1 to several hours daily) elevations in heart rate, blood pressure, cardiac output, and cardiac chamber volumes.9 Heavy and sustained ET generates large quantities of free radicals55 that likely outstrip the buffering capacity of the system, leaving these individuals susceptible to oxidative stress and transient cardiomyocyte dysfunction.45 This repetitive cycle may stimulate immune cells, including lymphocytes, macrophages, and mast cells, to secrete cytokines that signal the myofibroblasts to proliferate and secrete procollagen, which is then cross-linked to form mature collagen,45 eventually resulting in fibrosis deposited in patches in the myocardium and more diffusely in the large arteries.9,46

**PROARRHYTHMIC EFFECTS OF EXCESSIVE ENDURANCE ET**

Although it has been recognized that elite-level endurance athletes commonly have electrocardiographic abnormalities and atrial and ventricular ectopy,28,44,54 these functional adaptations traditionally have not been thought to predispose to serious arrhythmias or SCD. However, it appears that adverse cardiac remodeling induced by excessive ET can create an arrhythmogenic substrate, and rhythm abnormalities may be the most common CV problems encountered by veteran endurance athletes.29,31,54 Indeed, long-term sustained vigorous aerobic ET such as marathon or ultramarathon running or professional cycling has been associated with as much as a 5-fold increase in the prevalence of atrial fibrillation.19,30,31,37,55-63

Potential mechanisms underlying the association of long-term excessive exercise and atrial fibrillation are speculative but may include increased vagal and sympathetic tone, bradycardia, inflammatory changes, atrial wall fibrosis, and increased atrial size.59 Some data indicate that atrial size may be larger in veteran endurance athletes than in age-matched sedentary controls.60 Indeed, the left atrium may be enlarged in as many as 20% of competitive athletes, and this may be a predictor for atrial fibrillation.59,64

In addition, complex ventricular ectopy, including ventricular tachycardia and rarely SCD,11 occurs even in very fit individuals.12-20 Despite the fact that these studies generally excluded athletes with findings suggestive of familial arrhythmogenic RV dysplasia, the VAs typically originate from a mildly dysfunctional RV and/or the interventricular
The patchy myocardial fibrosis (fibrillary collagen deposition) that may develop as a reparative response to damaged myocardium can favor reentry, which is well established as a substrate for arrhythmia.29,51

Long-term extreme endurance ET and competition also stimulate multiple other disruptions, including episodic release of excessive catecholamines with resultant coronary vasoconstriction, chronic elevations of heart rate, changes in free fatty acid metabolism, lactic acidosis, and metabolic derangements.41

RISK STRATIFICATION FOR ENDURANCE ATHLETES
Currently, we have no proven screening methods for detecting potential CV pathologic changes associated with extreme endurance ET. A logical strategy for now might be to deploy postcompetition cardiac biomarkers, echocardiography, and/or advanced imaging such as cardiac MRI to identify individuals at risk for and/or with subclinical adverse structural remodeling and substrate for arrhythmias, but the cost would likely be prohibitive.65 Computed tomography for CAC scoring may be useful, particularly for those older than 50 years who have been training extensively for and competing in extreme endurance events. Exercise testing generally has not been found to be helpful in screening extreme endurance athletes, nor has cost-effectiveness or clinical yield been found with the other testing described earlier.

An obligatory pattern of compulsive and excessive daily exercise has been described that may have adverse long-term mental and physical health consequences.67 A questionnaire developed to identify obligatory exercisers may be useful for screening veteran endurance athletes.68

CONCLUSION
In some individuals, long-term excessive endurance ET may cause adverse structural and electrical cardiac remodeling, including fibrosis and stiffening of the atria, RV, and large arteries. This theoretically might provide a substrate for atrial and ventricular arrhythmias and increase CV risk. Further investigation is warranted to identify the exercise threshold for potential toxicity, screening for at-risk individuals, and ideal ET regimens for optimizing CV health. For now, on the basis of animal and human data, CV benefits of vigorous aerobic ET appear to accrue in a dose-dependent fashion up to about 1 hour daily, beyond which further exertion produces diminishing returns and may even cause adverse CV effects in some individuals.

Consensus Guidelines for Physical Activity and Public Health from the American Heart Association

FIGURE 6. Delayed gadolinium enhancement in 5 athletes. Images of 5 athletes in whom focal delayed gadolinium enhancement was identified in the interventricular septum (arrows), compared with a normal study in an athlete (top left). From *Eur Heart J.*,53 with permission.

FIGURE 7. Proposed pathogenesis of cardiomyopathy in endurance athletes. BNP = B-type natriuretic peptide; CK-MB = creatine kinase MB; LV = left ventricle; RA = right atrium; RV = right ventricle; SCD = sudden cardiac death.
and American College of Sports Medicine call for at least 150 minutes per week of moderate ET or 75 minutes per week of vigorous ET in the general adult population. Those guidelines also suggest that larger doses of ET may be necessary in some groups, such as those with or at risk for CHD (30 to 60 minutes daily), adults trying to prevent the transition to overweight or obesity (45 to 60 minutes per day), and formerly obese individuals trying to prevent weight regain (60 to 90 minutes per day). The guidelines also caution that high-intensity ET increases risk of musculoskeletal injuries and adverse CV events.1

Abbreviations and Acronyms: CAC = coronary artery calcium; CHD = coronary heart disease; CV = cardiovascular; EF = ejection fraction; ET = exercise training; LV = left ventricular; MRI = magnetic resonance imaging; PA = physical activity; RA = right atrium; RV = right ventricular; SCD = sudden cardiac death; VA = ventricular arrhythmia

Correspondence: Address to James H. O’Keefe, MD, 4330 Wornall Rd, Ste 2000, Kansas City, MO 64111 (jokeefe@saint-lukes.org).

REFERENCES


