Review Article

Prevention of sudden cardiac death in athletes, sportspersons and marathoners in India

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A B S T R A C T

The annual incidence of sudden cardiac death (SCD) in athletes is significantly lower than the general population. However, when SCD occurs in an athlete during sporting event or training, it sends shockwaves in the society and raises questions about cardiovascular effects of sports and exercise. This document reviews the causes and mechanism of SCD in sports and exercise in young and older athletes. In the Indian context, we suggest 'pre-participation screening' of young and older athletes and consider a 'supervised, graded exercise regime' for the uninitiated, older sports participant. Finally, the document proposes medical infrastructure required to successfully revive a victim of sudden cardiac arrest during a sporting event.

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Abbreviations: ACLS, advanced cardiac life support; AED, automated external defibrillator; ALCA, anomalous left coronary artery; ARCA, anomalous right coronary artery; ARVC, arrhythmogenic right ventricular cardiomyopathy; ASCVD, atherosclerotic cardiovascular disease; BCLS, basic cardiac life support; CAD, coronary artery disease; CI, confidence interval; CMR, cardiac magnetic resonance; CPR, cardiopulmonary resuscitation; CV, cardiovascular; CPVT, catecholaminergic polymorphic ventricular tachycardia; EMR, emergency medical response; HCM, hypertrophic cardiomyopathy; IDCM, idiopathic dilated cardiomyopathy; LGE, late gadolinium enhancement; LV, left ventricle; LVH, left ventricular hypertrophy; LVWT, left ventricular wall thickness; MI, myocardial infarction; RR, relative risk; RV, right ventricle; SCA, sudden cardiac arrest; SCD, sudden cardiac death; STEMI, ST elevation myocardial infarction; VT, ventricular tachycardia; VF, ventricular fibrillation; WPW, Wolff-Parkinson-White.

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### 1. Introduction

It is widely known and proven consistently by many studies that regular physical exercise significantly reduces the risk of cardiovascular death including SCD compared to sedentary lifestyle.\(^1,2\) However, exercise and sports-related physical activity sometimes result in SCD especially in those athletes with underlying silent or unknown coronary, myocardial or valvular heart disease and channelopathies. This suggests that in predisposed individuals exercise may act as a trigger for the development of malignant arrhythmias. The occurrence of such an event in an athlete especially during a sporting event draws a lot of media attention and often raises doubt about the cardiovascular benefits of exercise.

A pre-participation screening consists of structured history and physical examination, and when necessary ECG and an echocardiogram to identify young, future athletes at risk of SCD. The older uninitiated individuals participating in popular marathons also need to be evaluated, especially for risk factors and symptoms of coronary artery disease (CAD). High-risk individuals may undergo exercise testing and be guided appropriately to primary prevention strategies; they likely will benefit from a supervised exercise program.

This document intends to briefly review the causes and mechanisms of SCD in athletes, sportsperson and marathoners; suggest pre-screening methods to be employed in India; recognize the physiological and pathological ECG and echocardiogram findings; recommend the manner of supervised exercise training and finally the resuscitative measures that should be available and implemented in the rare occurrence of such an event.

This guidance will benefit trainers and organizers of such events, sports physicians and additionally educate the participants. It hopefully will promote governmental sports and health ministry to take necessary steps to prevent SCD in both competitive and leisure athletes and in particular marathon runners, an ever-expanding sporting activity. In India, there is a need for national registries of athletic/sports/marathoners SCD rates and collection of data on pre-participation screening. This should help to frame customized guidelines to prevent SCD in athletes, sportspersons and marathoners.

### 2. Definition of an athlete

American College of Cardiology Sports and Exercise Cardiology Think Tank defines **athlete** as any individual who engages in routine vigorous physical exercise in the settings of competition, recreation, or occupation.\(^3\)

A **competitive athlete** is defined as an individual . . .

1. Who participates in an organized team or individual sport,
2. Who engages in regular competition against others,
3. Who places a high priority or premium on athletic excellence and achievement,
4. Who engages in vigorous training in a systematic fashion to achieve all the above goals.\(^3,7\)

### 3. Epidemiology and causes of SCD in sports and exercise

The annual incidence of sports-related SCDs was reported to be 4.6 per 1 million population in a five-year prospective study\(^12\) as compared to 50–100 per million in the general population. The incidence and underlying cause of sudden death varies depending on the age, gender, race, ethnicity of the athlete, training level of athlete as well as the type of sport. Most sports-related SCD occurs in the middle-age group of 35 years or older\(^13\) or recreational athletes\(^14\) and 80% of these deaths are due to atherosclerotic CAD. The risk of sudden cardiac arrest (SCA) or SCD during training for the sports event is approximately 5-fold greater.\(^12\) All adverse event rates are higher in occasional runners (1 per 7500–18000) compared with trained marathon runners (1 per 50,000–200,000).\(^15\)

In younger athletes, the non-medical and traumatic causes constitute more than half of all deaths.\(^16\) The prevalence of cardiovascular (CV) diseases that predispose to SCD during sports in young athletes is estimated to be 0.2 to 0.7%.\(^4,7\) The most frequent cardiac causes of SCD in young athletes are Sudden Arrhythmic Death Syndrome (56%), hypertrophic cardiomyopathy (HCM) (36–48%) and congenital anomalous coronary artery origin from opposite sinuses (14–17%) while the less frequent causes are arrhythmogenic right ventricular cardiomyopathy (ARVC) (4–11%), myocarditis (6–7%) and ion channelopathies (4%).\(^10,11,17,18\) Atherosclerotic CAD is responsible for SCD in only 2–3% of younger athletes. The reported incidence of SCD in young athletes is variable in the literature. Maron et al\(^17\) reported 0.6 SCD per 100,000 person-years in young competitive athletes in United States. Van Camp et al\(^19\) reported 0.4 SCD in 100,000 athletes per year, in high school and college athletes. Among young Italian athletes, Corrado et al\(^10,11\) reported the incidence of SCD 3.6/100,000 person-years before routine pre-participation screening and 0.4/100,000 person-years after starting routine pre-participation screening. Among American Collegiate Athletes, Harmon et al\(^16,20\) reported the overall incidence of SCD 1:53703 athlete-
years. The SCD incidence was higher in Black athletes (1:21491 athlete-years) compared to White athletes (1:68354 athlete-years). The SCD incidence was highest in black basketball athletes (1 in 5348 athlete-years).

Various epidemiological studies on athletes and marathon runners published in the last decade have helped to establish the incidence and causes of SCD.21–28 The causes of SCD in sports-persons are listed in Table 1.

4. Mechanism of SCD during sports and exercise

The catecholamine surge during exercise interacts unfavorably with the underlying arrhythmogenic substrate. Intense exercise can cause dehydration, hyperventilation, electrolyte imbalance and increased platelet aggregation, further facilitating ventricular tachycardia or fibrillation.21

4.1. Younger athletes

As mentioned earlier, HCM, idiopathic left ventricular hypertrophy (LVH), coronary anomalies, myocarditis, ARVC, Marfan’s syndrome, valvular heart disease, or ion channelopathies form an underlying substrate in the younger athlete.7,9,21 Athletic training itself may unfavorably alter the substrate in HCM, ARVC and Marfan syndrome.28 The sports involving start-stop jolts like basketball and soccer have highest incidence of SCD in younger athletes than the endurance events like running.7,16,21,30

HCM: the myocardial disarray and interstitial fibrosis and microvascular ischemia form a substrate for ventricular tachycardia (VT) or fibrillation (VF).

ARVC: the progressive fibrofatty replacement of the right ventricular (RV) myocardium forms the substrate for the arrhythmia. The propensity to ventricular arrhythmias with sport is more than at rest.10

Congenital coronary anatomy anomalies: the anomalies causing SCD during sports are left coronary artery origin in the right sinus of Valsalva (ALCA) and right coronary artery origin in the left sinus of Valsalva (ARCA).9 The ALCA or ARCA may have acute-angled take off or slit like ostial stenotic opening or intramural course. The proximal segment of the vessel may get compressed as it courses between the pulmonary artery and ascending aorta. The mechanism of SCD is related to exercise induced acute ischemia in left or the right coronary artery territory.

Marfan syndrome: the tunic media of the aorta may undergo cystic medial necrosis causing aortic root dilatation. The increase in aortic pressure during exercise may cause aortic dissection or rupture.21

Wolff-Parkinson-White syndrome: is characterized by pre-excitation due to antegrade conduction via accessory atrioventricular pathway. Exercise-related atrial fibrillation may cause rapid antegrade conduction via the accessory pathway and result in VF and SCD.31

Congenital Long QT Syndromes: abnormal cardiac repolarization (QTc interval >470 ms in males and >480 ms in females) predisposes to polymorphic VT/VF during exercise. SCD occurs more commonly during swimming or diving due to adrenergic surge with sudden cold-water immersion.32

Brugada Syndrome: SCD is caused by polymorphic VT/VF, that is more frequent at rest. In athletes with Brugada syndrome, the increased parasympathetic tone and core body temperatures >40 °C may precipitate fatal arrhythmias.21

Catecholaminergic Polymorphic VT: exercise provokes multifocal ventricular premature beats or VT with beat to beat alternation of the frontal plane QRS axis (bidirectional VT).

Commotio Cordis: a single blow on the precordium strategically timed 10 to 30 ms before the T-wave peak on the electrocardiogram can induce VF and cardiac arrest in persons with normal heart.21 The Commotio Cordis related SCD is seen in contact sports involving player collisions (football, martial arts etc.) or sports involving projectile objects. Since it occurs in normal heart, it can’t be prevented by screening but can only be treated by prompt resuscitation measures on the field.

Table 1
Causes of SCD in sports.7,9,16,20,21

| Types                        | Subtype                          | Pathology                          |
|------------------------------|----------------------------------|------------------------------------|
| Structural Disorders         | Cardiomyopathy                   | HCM                               |
|                              | Marfan                           | Idiopathic LVH                     |
|                              | Valvular disease                 | ARVC                              |
|                              |                                  | MVP                               |
|                              |                                  | Aortic root dilatation/rupture/dissection |
| Disorders of Coronary Circulation | Congenital                  | ALCA from right sinus              |
|                              | Acquired                         | ARCA from left sinus               |
|                              | Ion channelopathies              | Atherosclerosis                    |
| Electrical disorders         | Ventricular Tachycardia          | Long QT syndrome                   |
|                              | Ventricular Fibrillation         | WPM syndrome                       |
|                              | Sports injury                    | Brugada syndrome                   |
|                              | Heat stroke                      | Short QT syndrome                  |
|                              | Infection                        | CPVT                              |
|                              | Performance enhancing drugs      | Sudden Arrhythmic Death Syndrome   |
|                              | Hyperthermia                     | Commmotio Cordis                   |
|                              |                                  | Physical trauma                    |
|                              |                                  | Ventricular fibrillation           |
|                              |                                  | Subacute Myocarditis               |
|                              |                                  | Myocardial infarction, Ventricular arrhythmias |
|                              |                                  | Ventricular arrhythmias            |

Abbreviations: – ALCA, Anomalous Left Coronary Artery; AR, Aortic Regurgitation; ARCA, Anomalous Right Coronary Artery; ARVC, Arrhythmogenic Right Ventricular Cardiomyopathy; AS, Aortic Stenosis; CPVT, Catecholaminergic Polymorphic Ventricular Tachycardia; HCM, hypertrophic cardiomyopathy; IDC, Idiopathic Dilated Cardiomyopathy; LVH, Left Ventricular Hypertrophy; MR, Mitral regurgitation; MVP, Mitral Valve Prolapse; SCD, Sudden Cardiac Death; WPM, Wolff-Parkinson-White
Myocarditis: immune mediated myocardial inflammation in viral myocarditis predisposes to VT/VF during exercise. The risk may continue up to 6 months in convalescent period.

Performance-enhancing drugs: These drugs often used illegally can act as trigger or lead to development of an arrhythmogenic substrate. Anabolic androgenic steroids cause premature atherosclerosis, hypertension (HT), dyslipidemia and myocardial infarction. Ephedrine use leads to HT and cardiomyopathy. Recombinant human erythropoietin can cause HT and thromboembolism. Human growth hormone use leads to myocardial hypertrophy and fibrosis.23,34

Premature coronary artery disease: Familial hypercholesterolemia in young adults causes premature coronary atherosclerosis and susceptibility to ischemia or myocardial infarction (MI).

4.2. Older athletes

In athletes older than 35 years of age, most deaths occur during long distance running sports.7,9 In RACE Paris registry, SCDs due to CAD were demonstrated to have acute thrombotic occlusion in all cases,15 however, the RACER registry of 10 million runners found that SCD in persons with CAD occurs due to fixed coronary stenosis without any thrombosis.26 An autopsy series of SCDS in general population showed higher incidence of plaque erosion (68% versus 23%) and intraplaque hemorrhage (71% versus 41%) in SCD with exercise versus SCD at rest.35

Therefore, there are two plausible hypotheses to explain the occurrence of ventricular tachyarrhythmia and cardiac arrest during sports activity –

1) Fixed coronary stenosis: the coronary flow cannot be augmented during exercise since the coronary vasodilatory reserve beyond stenosis is nearly exhausted and the diastolic period is shortened due to tachycardia. During exercise there is increased myocardial oxygen demand. The ischemia may get exacerbated by abrupt cessation of exercise that leads to reduced venous return, fall in blood pressure in a vasodilated state resulting in coronary hypoperfusion. The acute ischemia is further worsened by electrolyte imbalance, heat stroke36,37 and excess circulating catecholamines.29 This eventually causes malignant ventricular arrhythmias either at peak effort or immediate post exercise.

2) Acute plaque rupture: The increased wall shear force on vulnerable plaque during exercise and catecholamine induced coronary spasm along with endothelial dysfunction cause erosion of the thin fibrous cap and/or intraplaque hemorrhage and thrombosis. The high levels of circulating catecholamine potenti- ated platelet aggregation. The acute ST segment elevation myocardial infarction (STEMI) or non-STEMI leads to malignant ventricular arrhythmias.29

4.3. Exercise paradox or sports paradox

The exercise induced hemodynamic stress and catecholamine surge increase the incidence of acute ischemia, myocardial infarction and SCD during and up to 1 h post-exercise compared to the sedentary hours. The incidence of SCD with exercise is 8–16 times higher38,39 and the incidence of MI with exercise is 6–10 times higher40,41 compared to during resting hours. This increased relative risk of SCD and acute MI associated with every bout of vigorous exercise is reduced dramatically, in a dose-dependent fashion in the long term, by habitual and regular exercise. This is known as exercise or sports paradox.

The moderate to high intensity exercise, performed at regular intervals over a long period of time, has positive pleiotropic effects42 in reducing the overall risk of atherosclerosis and acute coronary syndrome. Compared to sedentary individuals, the relative risk of SCD during ‘exercise hours’ is reduced by 7–10 folds and the relative risk of acute MI during ‘exercise hours’ is reduced by 50 folds in individuals performing regular vigorous exercise.12 The intensity and frequency of regular exercise has inverse dose response relationship with relative risk of MI and SCD during exercise bout. Therefore, an active individual will have fifty-fold lower risk of MI during the exercise hour and will have up to two-fold lower risk of MI during the remaining sedentary hours of the day, showing the net benefit of exercise.43

5. Pre-participation screening

Sudden cardiac death in apparently fit appearing athletes mostly occurs in the absence of any warning symptoms or history of heart disease. Pre-participation screening to identify the presence of silent heart diseases can reduce the risk of sports related sudden death.

5.1. Young athletes

History, physical examination and ECG are the least expensive means of pre-participation screening and constitute the primary screening tool in most guidelines.7

5.1.1. History and examination

The current AHA recommendations for pre-participation screening of a young, competitive athlete includes essential elements in history and physical examination.7 This should be undertaken by the athlete’s personal/family physician upon entry into competitive sports at the school or college levels. The physical examination should include careful auscultation (in various positions and with maneuvers if needed) to detect any heart murmur especially a dynamic murmur due to left ventricular outflow tract obstruction. If there is any suggestion or suspicion of any heart disease based on this evaluation, further testing and evaluation by a cardiologist is recommended. The young athlete with family history of SCD or known cardiomyopathy should be evaluated with more elaborate history and clinical examination. This group of athletes should be referred to cardiologists and mandatorily have ECG and echocardiographic evaluation at baseline and at intervals. Further imaging and electrophysiological testing can be planned at the discretion of the cardiologist.

5.1.2. ECG screening

The addition of 12-lead ECG to history and physical examination can increase the sensitivity of pre-participation screening in detecting significant cardiac abnormalities.44,45 ECG screening can identify cardiac diseases that manifest with specific findings such as ion channelopathies (for e.g., Long QT syndrome, Brugada syndrome, WPW syndrome, HCM and ARVC. However, non-specific ECG changes may be present in many individuals who are already performing athletic activities and may create confusion in the diagnosis.46,47 Pre-participation screening of young athletes with ECG, in Veneto region of Italy, showed reduction of incidence of SCD from 3.6/100,000 person-years to 0.4/100,000 person-years (p < 0.001), representing 90% reduction in sports related mortality.41 Most of the reduced incidence of SCD was achieved by detecting young athletes with cardiomyopathies like HCM and ARVC leading to their disqualification from competition. However, in a study from Israel, pre-participation screening of athletes with ECG and stress test was not associated with reduction in adverse cardiac events. The average incidence in the decade, before and after the mandated screening, was 2.54 and 2.66 events per 100,000 person-years, respectively.48 It is estimated that 30% of lethal CV causes like congenital coronary anomalies, 10–30% cases of cardiomyopathies (HCM/ARVC), aortic dilatation, IDCM and premature CAD cannot be identified by pre-participation screening.
with ECG. There is a concern about sensitivity, specificity, practicality, and cost-effectiveness of ECG screening strategy for the conditions (HCM, ARVC, ion-channelopathies) that have low prevalence in the young population. There is a divide in North America versus Europe guidelines regarding incorporating ECG as a mandatory test for pre-participation screening. Importantly, most of the rich sports associations have made ECG mandatory for screening prospective athletes. In Indian ECG is a relatively inexpensive test, and therefore suggested for all athletes/sports-person and marathoni. However, understanding the limitations of ECG is paramount to avoid unnecessarily barring athletes from pursuing their career of choice because of physiological changes in the ECG. At the same time, it should be borne in mind that a normal ECG does not rule out presence of heart diseases. Table 2 outlines the proposed pre-participation screening for young athletes in India.

5.2. Older athletes

A stepwise screening process as suggested by Chugh et al. is likely to be useful in detection of occult heart disease in the older athlete. It essentially involves a detailed history including that of coronary risk factors, physical examination and an ECG in all subjects as the first step. Any suspicion of heart disease or a high risk of CAD will necessitate evaluation by a cardiologist as a part of second step to assess the need for further tests like stress test, echocardiogram, other imaging tests or cardiac catheterization (Table 3).

5.2.1. Exercise ECG testing

The detection of silent coronary artery disease is one of the goals of pre-participation evaluation of athletes especially above 35 years of age. Exercise ECG testing is often performed as a part of screening in asymptomatic athletes and for evaluation of symptoms suggestive of CAD. The wide availability, low cost and known diagnostic value for detection of critical coronary stenosis in individuals at intermediate risk, make stress test useful in evaluating older athletes. However, the diagnostic accuracy of exercise testing depends on the pre-test probability of the disease and its prevalence in the community. Hence, its usefulness in asymptomatic individuals without coronary risk factors is limited. On the other hand, it can be useful for detection of CAD in those athletes who have coronary risk factors with an estimated 10-year-risk of CAD in the intermediate category. Although, an abnormal stress test indicates the presence of significant stenosis of one or more coronary arteries; a stress test cannot detect a non-significant stenosis due to a vulnerable plaque. Since SCD due to acute coronary syndrome can also occur because of rupture of a non-significant vulnerable plaque, the value of exercise ECG testing for predicting occurrence of sudden cardiovascular events during exercise is limited.

Many guidelines have incorporated the use of exercise ECG testing as a pre-participation screening in individuals at higher risk of CAD based on consensus, however scientific data is lacking.

6. ECG interpretation in athletes

The role of ECG for screening in athletes gets further compounded by the prevalence of abnormal findings on the ECG that occur as a result of athletic training itself in the absence of any heart disease. These “abnormal” ECG manifestations occur either due to increased vagal tone or due to increase in muscle mass and

Table 2
Pre-participation screening in young, competitive athletes.

| History | 1. Unexplained or exertional syncope  
2. Exertional chest pain/dyspnea  
| Family History | 1. Premature SCD (<50 years and in first degree relative)  
2. Known h/o hypertrophic or dilated cardiomyopathy, Long QT syndrome, CPVT, ARVC, Brugada, Marfan syndrome  
| Examination | 1. Blood pressure — both upper and lower limbs  
2. Heart murmur cardiomegaly  
3. Marfanoid features  
4. Xanthelasma  
| ECG | 1. Ischemia/infarction  
2. Chamber enlargement  
3. Long QT/WPW/ARVC/Brugada  
Abbreviations: ARVC, Arrhythmogenic Right Ventricular Cardiomyopathy; CPVT, catecholaminergic polymorphic ventricular tachycardia; SCD, Sudden Cardiac Death; WPW, Wolff-Parkinson-White.

Table 3
Steps in pre-participation screening for older athletes.

| Step | History | Examination | ECG |
|------|--------|-------------|-----|
| 1:   | Exertional chest pain, Syncope, Palpitations  
Family history of MI, SCD and Coronary Risk Factors  
Coronary Risk factors — HT, DM, dyslipidemia, smoking, family history of premature CAD  
Murmur, abnormal heart sound/s, evidence of cardiomegaly  
Ischemia, infarction, chamber enlargement or channelopathy  
If Step 1 positive → Fit for exercise training  
If Step 1 negative → Evaluate Step 2  
Step 2: Assessment by cardiologist for necessity of further tests | Stress test  
Echocardiography  
Cardiac MRI  
Cardiac Catheterization |
Abbreviations: CAD, Coronary Artery Disease; DM, Diabetes Mellitus; HT, Hypertension; MI, Myocardial Infarction; MRI, Magnetic Resonance Imaging; SCD, Sudden Cardiac Death.
cardiac size. Their occurrence is also influenced by age, sex, race & ethnicity and most importantly by the type of sport and duration of athletic training. Athletes involved in endurance sports like long-distance running are at maximum chance of training-related ECG manifestations.52

The ESC has provided recommendations for ECG interpretation in athletes and have grouped them into those resulting from physiological adaptation of cardiac autonomic nervous system and those that are likely due to underlying heart disease. The physiological adaptation related ECG abnormalities are seen in up to 80% of athletes whereas ECG changes suggesting heart disease are seen in less than 5% and these warrant further cardiac evaluation.57 A summit on “ECG interpretation in athletes” held in Seattle, Washington USA in 2012 involving leading cardiologists from various international societies laid down criteria that if present on ECG indicate possible underlying heart disease requiring further diagnostic evaluation.53

More recently, an international consensus for ECG interpretation in athletes was published that provides expert opinion-based recommendations linking specific ECG abnormalities and the secondary evaluation for conditions associated with SCD.54 The guidelines classified the ECG findings as normal, borderline and abnormal and suggested that further evaluation is required in the presence of abnormal findings or two or more borderline findings. In the absence of cardiac symptoms or family history, presence of normal findings or borderline changes in isolation do not warrant further evaluation. (Fig. 1)

7. Athletes heart and its differentiation from pathological states

The electrical, structural and functional cardiac adaptation observed in the athletes is termed as athlete’s heart. The clinician should differentiate these physiological adaptations from pathological changes using clinical, ECG, imaging data and objective functional capacity.

7.1. Athletes heart versus HCM

Athletes show 10–20% increase in left ventricular (LV) wall thickness (LVWT). LVWT of 13–16 mm falls into a ‘grey zone’ and more clinical and imaging data is required to differentiate athlete’s heart from HCM. Echocardiography showing a small LV size, enlarged LA, asymmetric or focal hypertrophy along with abnormal diastolic relaxation suggests HCM. Non-sustained VT on Holter, CMR for LGE and genetic testing further help to establish the diagnosis of HCM.

7.2. Athletes heart versus ARVC

The overall cardiac chamber enlargement in an athlete might be confused with RV dilatation seen in ARVC. Their differentiation is extremely important since exercise not only predisposes a patient with ARVC to SCD, but helps in the progression of the disease and unfavorably modifies the substrate. The RV enlargement compatible with ARVC diagnosis is observed in 22% of black athletes and 29% of white athletes.55–57 A positive family history along with ECG and echocardiographic findings may help to identify ARVC and typical findings on CMR imaging confirms the diagnosis.

7.3. Athletes heart versus IDCM

Athletes show 10–15% increase in both left and right ventricular cavity size. In Olympic athletes, 45% have LV cavity size more than upper limits of normal and 14% have LV cavity more than 60 mm.54 Family history and additional ECG changes (LBBB, T wave inversion in infero-lateral leads) help to establish the diagnosis of IDCM. Diastolic dysfunction and failure of improvement in LV systolic function on exercise echocardiography suggests IDCM. Additionally, low peak VO2 on cardiopulmonary testing, non-sustained VT on Holter, LGE on CMR and positive genetic testing favor a diagnosis of IDCM.
8. Exercise training

Prior to beginning a program of vigorous exercise, usually defined as >6 METS (metabolic equivalents), it is recommended to undergo medical clearance by a physician, familiar with sports medicine. Evaluation should be directed to screen for the presence of cardiovascular, pulmonary and metabolic diseases and their risk factors. This is particularly important for those who do not regularly participate in exercise or sports activities.18

Training plans for full marathons should be at least 16–20 weeks long, after having a baseline ability to run at least 8 km, several times a week. Training plans usually include running distances of 5–10 km, 3–5 days of the week. One day a week, athletes should run for a longer distance (upward of 10 km), building the distances up progressively every week. It is important to increase the weekly mileage in a gradual manner, to lessen the chances of injury; with a rule of thumb being, an increase of not more than ten percent. Athletes should train with optimal individualized dose and intensity19 of exercise or activity under guided supervision. Individuals more than 35 year should undergo supervised or guided exercise training programs with gradually increasing intensity. Also, the program should religiously include ‘warm up’ and ‘cool down’ exercises before and after every bout of vigorous exercise respectively. Training heart rate, should generally be in the range of 60–90% of maximal heart rate (obtained on a symptom-limited graded exercise test, or age-predicted). Participants should acclimatize to the venue climate before the race and should not participate if suffering from fever or respiratory tract infection to prevent heat stroke. They should be made aware of common symptoms of cardiovascular insufficiency, such as chest discomfort and unusual shortness of breath with exertion. Since marathon running is relatively new in India, it is advisable for participants to train under the guidance of a qualified coach.

9. Medical facilities during sporting events

Planning the delivery of medical services at an event should include a risk assessment and buy-in from multiple agencies to ensure that first-aid and medical services are provided for, a thorough communications plan is in place, emergency access/exit routes are mapped out, and staff & volunteers are trained in escalation and evacuation procedures.60 There should be a unified command centre for co-ordinated monitoring and communication of the event. Security approvals should be taken from the concerned government agencies and also have them part of the planning process.

Medical facilities during extreme endurance sporting events are usually made available to take care of the common risks like dehydration, heat exhaustion & sporting injuries. However, cardiovascular events, including cardiac arrest, although uncommon, are of prime concern due to their life-threatening nature, facilities to diagnose and manage these at sporting event venue itself should always be available. These life-threatening events are more frequently observed to occur during the end of the race suggesting a sudden surge in acceleration to reach the finish line setting off a chain of bodily reactions leading to cardiac arrest.15,61

While planning for medical provisions for an event, factors such as event type, weather (especially heat index), presence of alcohol and/or illicit drugs, number of participants, duration of the event, crowd demographics and venue characteristics should be taken into consideration. Appropriate judgement of the medical problem presentation rate and ambulance transfer time are very useful in understanding the medical needs of a sporting event.62

9.1. Staffing of the event

Number of staff at an event would be determined by the number of participants and the extent of medical coverage required. A marathon requires a mix of facilities with basic medical aid stations and ambulance evacuation. The event should have a medical director, who should preferably be a sports medicine physician, sports cardiologist, cardiologist or general physician with experience of planning mass sporting events. At every medical station, along the route there should be one physical therapist, one nurse and emergency medicine paramedic, with BCLS training and equipment to provide first aid. Majority of medical staff should be diverted toward the finish line as the race moves onwards. There should be at least one medical professional per 100 runners, in events with running distances longer than 10 km.

9.2. Physical infrastructure

Infrastructure to facilitate rapid diagnosis, treatment, and triage to appropriate centres, is essential for optimal care of participants in such mass gatherings. Medical services at mass sporting events can be delivered on foot, on bicycle, on motorbikes, on golf carts or other motorized vehicles, ambulance, first-aid stations and local hospitals. There should be a central medical station set up at the finish line which should be equipped like a mini ICU. There should be one aid station every 3 km of the route. Routes which are out-and-back (loop), make it easier to cover, since runners on both sides of the road can avail the facilities and only ‘half the number of physical structures’ and personnel are required. Volunteers should be stationed as ‘spotter’ who have been trained to identify athletes in distress and who are aware of the medical escalation protocol. Ambulances should be placed at regular intervals along the course of the race, and especially near the finish line and halfway between the first-aid stations.

9.3. Communication

The unified command centre is the main hub to monitor and coordinate all activities related to the event. There should be designated staff for coordinating medical emergencies. Deployed mobile units and staff at first-aid tents should be in touch with one another through mobile phones and walkie-talkies. The latter is preferred as the entire medical team can listen and respond at once. The route map with various facilities should be with every person involved in organizing the event. The chain of command, flowcharts for identification & referral and activating emergency medical response should not only be part of the training module but should also be documented and be part of the essential docket for the event.

9.4. Handling an emergency

First responders should be trained to identify symptoms, trigger the emergency medical system in place, perform Cardio-Pulmonary Resuscitation (CPR), and use AEDs.52,63 First responders should be able to summon emergency medical units and/or ambulances based on symptoms. The medical aid units should be able to identify and treat hyperthermia. Hyponatremia is a very real and under-recognized life-threatening situation in races.64 Protocols for the assessment of dehydration, asthma, chest pain, syncope, or exercise-induced cramping should be part of the training module as well.55 Local hospitals should be part of the planning process in order to be prepared for life threatening situations at the day of the event, and should be alerted about receiving casualties during the event.
The survival in sports related SCD doubles with active CPR and rapid defibrillation. 14,15,16,6 The SCDs in US Marathons was 1 per 55,000 finishers before 1995, which reduced to 1 per 220,000 finishers during 1995 to 2004 as a result of emergency medical response (EMR) facilities. To be effective, the EMR team should conduct regular emergency drills and CPR practice and should have “hotline” to summon emergency medical transportation services. EMR should review the quality metrics like time to CPR/defibrillation and the outcome at least annually.

Communities including athletes, coaches, fitness trainers and spectators, should be trained in recognition of SCA and basic CPR for increasing the chances of immediate bystander CPR during sporting events. “Do Not Start marathon” temperature and air pollution thresholds should be decided to reduce heat-stroke risks during running events in hot and humid places.

10. Conclusion

Sports related SCD could be prevented and minimized by understanding the causes and mechanisms of such events. In the younger athlete, pre-participation screening with history (including family history), physical examination and ECG should help identify those at high risk for SCD and a detailed examination by a cardiologist can be considered for those suspected of heart disease or risk on initial screening. The older athlete is more likely to suffer SCD than the younger athlete; and since CAD is the most frequent cause for such events, their pre-participation screening should include evaluation for coronary risk factors and when necessary a stress test. Importantly, preventive strategies should be advised in older athletes with coronary risk factors. Supervised and graded exercise training is likely to prevent coronary events. It needs to be emphasized that regular exercise remarkably reduces the risk of MI and SCD.

Unfortunately, despite the best screening programs and exercise regimes, rare occurrence of SCD during athletic events is a reality. Appropriate planning and co-ordination of sporting events by the various agencies, especially the medical team and the paramedics, should help to identify and revive an athlete with sudden cardiac arrest.

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