Introduction

Regular exercise has many favorable effects on the cardiovascular system. However, intense physical activity, especially in a competitive fashion, may become hazardous to genetically vulnerable individuals.

A competitive athlete is one who participates in an organized team or individual sport that requires regular competition against others as a central component, places a high premium on excellence and achievement, and requires some form of systematic (and usually intense) training. Soccer, basketball, running, cycling, and swimming are some examples of competitive sports with high dynamic activity, whereas gymnastics, weight lifting, and sailing are examples of competitive sports with high static activity.

Sudden cardiac death (SCD) in sports and exercise is defined as cardiac arrest occurring during or within 1 hour of exercise or sports-related activity, and genetically transmitted cardiovascular diseases are among the most common pathologies leading to SCD in competitive athletes (1).

Sudden death of a competitive athlete has a mass and undesirable effect on the community. Properly diagnosing inherited cardiovascular pathologies, making appropriate recommendations on physical activity in athletic population, and providing close supervision to diseased athletes are the main duties of cardiovascular specialists.

The aim of our document is to summarize the various aspects of the most common genetic cardiovascular diseases in athletes and to make recommendations for diseased athletes and athletes with implanted cardiac pacemakers/Implantable Cardioverter Defibrillators (ICDs) based on the current literature. Our general recommendations are summarized in Figure 1 and discussed below in detail.

The incidence of SCD in athletic population varies between 0.5 and 13 deaths per 100,000 athletes according to current literature (2, 3).

The most common genetic cardiovascular diseases which leads to SCD varies among different regions of the world. In the United States (US), hypertrophic cardiomyopathy (HCM) is responsible for one-third of the mortality in the athletic population.
and congenital coronary anomalies is the second in frequency (4). In the Veneto region of Italy, arrhythmogenic right ventricular cardiomyopathy (ARVC) has been reported as the most common cause of SCD in young athletes (5). Another study which investigated the causative pathologies of SCD in athletes with detailed post mortem evaluation found that idiopathic left ventricular hypertrophy/fibrosis and ARVC were the most common causes of SCD in athletes (6). The etiology of SCD in competitive athletes involves a wide range of pathologies as listed in Table 1 and discussed below.

**Cardiomyopathies**

**Hypertrophic cardiomyopathy**

HCM is an inherited disorder defined by the presence of increased left ventricular (LV) wall thickness that cannot be explained by abnormal loading conditions of a cardiac or a systemic disease. LV wall thickness of ≥15 mm in at least one segment measured by an imaging technique is required for the definite diagnosis of HCM (7). Since HCM has been accepted as one of the major pathologies that leads to SCD in young athletes (4, 8), it is of paramount importance to detect the disease before athletic participation to make the distinction between HCM and physiological hypertrophy and manage patients during the sporting activity.

Main abnormalities on electrocardiography (ECG) of athletes with HCM are large QRS voltages, T wave inversion, ST depression, and pathological Q waves. On echocardiographic examination, most athletes with HCM had larger LV cavity dimensions with lesser LV hypertrophy and better indices of diastolic function compared to non-athletic HCM patients. The pattern of LV hypertrophy in athletes with HCM is mostly asymmetric and focal in contrast to the symmetrical pattern of physiological LV hypertrophy of the athlete’s heart. Apical hypertrophy is also seen in a substantial number of athletes with HCM. Healthy athletes show 10%–20% increase in LV wall thickness. LV wall thickness of 13–16 mm falls into a “gray zone”, and more clinical and imag-
Table 1. Etiology of sudden cardiac death in competitive athletes

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<td>o Non-compaction cardiomyopathy</td>
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<td>o Dilated cardiomyopathy</td>
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<td>o Brugada syndrome</td>
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<td>o Catecholaminergic polymorphic ventricular tachycardia</td>
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<tr>
<td>o Early repolarization syndrome</td>
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<tr>
<td>• Wolf Parkinson White syndrome</td>
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<tr>
<td>• Idiopathic left ventricular hypertrophy/fibrosis</td>
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<tr>
<td>• Coronary atherosclerosis</td>
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Patients with HCM are advised to restrict participation in athletic activities according to the AHA/ACC task force report which was published in 2015 (10). The latest ESC report stated that participation in intensive exercise programs and competitive sport should be considered on an individual basis. After full evaluation of the disease characteristics and risk determinants, patients with HCM who have high-risk clinical characteristics must be withheld from athletic activities (15). It is of paramount importance to discriminate the ones who have high-risk features of HCM when restricting athletes from competitive activities.

Non-compaction cardiomyopathy

Left ventricular non-compaction (LVNC) is a genetic cardiomyopathy which is characterized by prominent myocardial trabeculations and deep intertrabecular recesses associated with LV dysfunction. Heart failure symptoms, syncope, systemic thromboembolism, and VT are the main clinical presentations of the patients with LVNC (16). The main symptom was found to be syncope without prodromal symptoms during activity in athletes with LVNC (17). Although LVNC is rarely seen in athletes and accounts for the minority of SCDs in athletic cohorts, differentiating the morphological alterations of LVNC from the adaptive changes of an athlete’s heart is an important issue. There is no specific ECG finding for LVNC. LV hypertrophy, repolarization abnormalities, and QT prolongation were found to be the most common abnormalities on ECG (18). Sustained VT can also be seen in athletes with LVNC (17). Several echocardiographic criteria were identified for making the diagnosis of LVNC. The main findings on echocardiography are ≥2 ratio of non-compacted/compacted layer, presence of deep intertrabecular recesses filling with ventricular blood and numerous trabeculations protruding from LV wall with reduced ejection fraction (<50%) (16).

A study which compared the echocardiographic results of elite athletes with normal population and LVNC patients found that 20% of the athlete group expressed an increased number of LV trabeculation and 10% of the athletes had fulfilled the conventional echocardiographic criteria of LVNC. Authors stated that this may not be the true incidence of the disease and more stringent criteria for the diagnostic consideration of LVNC are required in this special population (19). According to another study, prominent LV trabeculation was found only in 1.4% of a large athlete population. Although 66% of the athletes who had LV hypertrabeculation match the echocardiographic criteria for LVNC, none of them had LV dysfunction, positive family history, pathologic cardiac MRI findings, and symptoms (20). Thus, athletes with hypertrabeculation have to be evaluated carefully and should not be diagnosed directly as LVNC. However, athletes with symptoms and/or high-risk clinical features should be restricted from the athletic activities.

Dilated cardiomyopathy

Dilated cardiomyopathy (DCM) is a myocardial disease characterized by dilated and hypokinetic LV with or without right ventricle dysfunction. Dilated cardiomyopathy may be idiopathic or originating from infection, inflammation, toxic agents or ischemia (21). Occasionally, DCM is a known cause of SCD in athletes (2, 3).

The ECG may be normal or exhibit similar changes to those of athletic individuals such as atrial dilatation, axis deviation or large QRS voltages, and T wave inversion in lateral leads.
### Consensus statements

<table>
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<tr>
<th>Hypertrophic cardiomyopathy</th>
<th>Recommendations</th>
<th>References</th>
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<tbody>
<tr>
<td>Athletes with HCM who are asymptomatic and do not have significant LVOT gradient could be supervised closely and may selectively participate in athletic activities.</td>
<td>🟠</td>
<td>10, 15</td>
</tr>
<tr>
<td>Athletes with HCM who have a history of aborted SCD, exercise-induced ventricular tachycardia, unexplained syncope, significant LVOT gradient, and abnormal blood pressure response to exercise have to be restricted from athletic activities.</td>
<td>🔴</td>
<td>10, 15</td>
</tr>
<tr>
<td>Genotype positive phenotype negative asymptomatic HCM patients without evidence of LV hypertrophy by imaging methods may participate in athletic activities.</td>
<td>🟠</td>
<td>10, 15</td>
</tr>
<tr>
<td>Genotype positive phenotype negative HCM patients should supervise closely to monitor the progression to hypertrophic phenotype.</td>
<td>🟠</td>
<td>10, 15</td>
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<tr>
<th>Non-compaction cardiomyopathy</th>
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<tbody>
<tr>
<td>Athletes who have a diagnosis of LVNC with normal EF, without symptoms and ventricular tachycardias on ambulatory monitoring and stress testing may not be restricted from athletic activities but close supervision needed.</td>
<td>🟠</td>
<td>15, 19, 20</td>
</tr>
<tr>
<td>Athletes with LVNC who have symptoms (especially syncope), reduced EF, thromboembolic events, and ventricular tachycardias on ambulatory monitoring or stress testing should be restricted from the athletic activities.</td>
<td>🔴</td>
<td>15, 19, 20</td>
</tr>
<tr>
<td>Asymptomatic athletes with hypertrabeculation and without a diagnosis of LVNC can participate in all competitive sports.</td>
<td>🟠</td>
<td>15, 19, 20</td>
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<tr>
<th>Dilated Cardiomyopathy</th>
<th>Recommendations</th>
<th>References</th>
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<tbody>
<tr>
<td>Asymptomatic athletes with DCM and mildly decreased LV systolic function (EF&gt; 40%), may selectively participate in athletic activities.</td>
<td>🟠</td>
<td>15, 22</td>
</tr>
<tr>
<td>In DCM, athletes with symptoms or reduced LV ejection fraction (&lt;40%) or frequent and complex ventricular tachyarrhythmia in ambulatory ECG monitoring or exercise tests or history of unexplained syncope should not be recommended to deal with athletic activities.</td>
<td>🔴</td>
<td>15, 22</td>
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<th>Myocarditis</th>
<th>Recommendations</th>
<th>References</th>
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<tbody>
<tr>
<td>If LV function and serum biomarkers of myocardial injury are normalized and no clinically relevant arrhythmia detected on 24 h ECG monitoring, it is reasonable for athletes with myocarditis to return athletic activities under close supervision after a healing period of 3 to 6 months.</td>
<td>🟠</td>
<td>10, 24</td>
</tr>
<tr>
<td>Athletes with myocarditis should be followed regularly in case of risk of recurrence and silent progression of the disease especially during the first 2 years.</td>
<td>🟠</td>
<td>10, 24</td>
</tr>
<tr>
<td>Athletes with myocarditis should be restricted from athletic activities for a period of 3 to 6 months.</td>
<td>🔴</td>
<td>10, 15, 24</td>
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<tr>
<th>Arrhythmogenic right ventricular cardiomyopathy</th>
<th>Recommendations</th>
<th>References</th>
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<tbody>
<tr>
<td>ARVC patients should not participate in high intensity athletic activities.</td>
<td>🔴</td>
<td>15, 27</td>
</tr>
<tr>
<td>Genotype positive phenotype negative ARVC patients should not participate in high intensity athletic activities.</td>
<td>🔴</td>
<td>15, 27</td>
</tr>
<tr>
<td>ICD implantation in an athlete with ARVC for the sole purpose of participation in high intensity athletic activity is not recommended.</td>
<td>🔴</td>
<td>15, 27</td>
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<tr>
<th>Commotio cordis</th>
<th>Recommendations</th>
<th>References</th>
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<tbody>
<tr>
<td>Commotio cordis survivors should undergo a complete cardiac study to exclude structural heart disease and underlying arrhythmic condition.</td>
<td>🟠</td>
<td>29, 30</td>
</tr>
<tr>
<td>After comprehensive evaluation of commotio cordis survivors, athletes without any underlying cardiac disease can safely return to athletic activities.</td>
<td>🟠</td>
<td>29, 30</td>
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</table>
Healthy athletes show a 10%–15% increase in both left and right ventricular cavity size. In Olympic athletes, 45% have LV cavity size over the upper limits of normal. Family history and additional ECG changes help establish the diagnosis of DCM. Diastolic dysfunction and failure of improvement in LV systolic function on exercise echocardiography suggests DCM. Additionally, low peak VO2 on cardiopulmonary testing, non-sustained VT on Holter ECG, LGE on cardiac MRI, and positive genetic testing favor a diagnosis of DCM (22).

### Consensus statements

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<th>Coronary artery anomalies</th>
<th>Recommendations</th>
<th>References</th>
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<tr>
<td>Athletes with anomalous origin of a coronary artery without either symptoms or positive stress test may be selectively participate in athletic activities after counseling with the athletes and/or parents of the athlete.</td>
<td><img src="https://via.placeholder.com/15" alt="Heart" /></td>
<td>15, 36, 43</td>
</tr>
<tr>
<td>After successful surgical repair; operated athletes with CAAs may consider to return athletic activities 3 months after surgery if the athlete is asymptomatic and a stress test shows no evidence of ischemia.</td>
<td><img src="https://via.placeholder.com/15" alt="Heart" /></td>
<td>15, 36, 43</td>
</tr>
<tr>
<td>Athletes with anomalous origin of a coronary artery which shows an interarterial course should be restricted from athletic activities before surgical repair.</td>
<td><img src="https://via.placeholder.com/15" alt="Heart" /></td>
<td>15, 36, 43</td>
</tr>
<tr>
<td>Athletes with anomalous origin of a coronary artery who exhibits symptoms or arrhythmias or signs of myocardial ischemia in stress tests should be restricted from athletic activities before surgical repair.</td>
<td><img src="https://via.placeholder.com/15" alt="Heart" /></td>
<td>15, 36, 43</td>
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<tr>
<th>Channelopathies</th>
<th>Recommendations</th>
<th>References</th>
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<tbody>
<tr>
<td>Athletes with a suspected cardiac channelopathy should be evaluated by an experienced heart rhythm specialist.</td>
<td><img src="https://via.placeholder.com/15" alt="Heart" /></td>
<td>46, 47, 52</td>
</tr>
<tr>
<td>It is advised to perform sports in places with on-board automated-external-defibrillator and near people who are already informed about the disease for athletes with channelopathy.</td>
<td><img src="https://via.placeholder.com/15" alt="Heart" /></td>
<td>46, 47, 52</td>
</tr>
<tr>
<td>Asymptomatic athletes with genotype positive phenotype negative channelopathy might be allowed to participate in all sports with appropriate precautionary measures.</td>
<td><img src="https://via.placeholder.com/15" alt="Heart" /></td>
<td>46, 47, 52</td>
</tr>
<tr>
<td>It is recommended that symptomatic athletes with any suspected or diagnosed channelopathy should be restricted from all competitive sports until a detailed evaluation has been completed, appropriate treatment has been applied and asymptomatic status on therapy has been provided for 3 months.</td>
<td><img src="https://via.placeholder.com/15" alt="Heart" /></td>
<td>46, 47, 52</td>
</tr>
<tr>
<td>Drugs which induce a Brugada-pattern on ECG and drugs which prolongs QT interval should be avoided in athletes with BrS and LQTS respectively.</td>
<td><img src="https://via.placeholder.com/15" alt="Heart" /></td>
<td>46, 47, 52</td>
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<tr>
<td>Dehydration, excessive sweating, electrolyte disturbances, and hyperthermia should be avoided for athletes with channelopathy.</td>
<td><img src="https://via.placeholder.com/15" alt="Heart" /></td>
<td>46, 47, 52</td>
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<tr>
<th>Wolff Parkinson White syndrome</th>
<th>Recommendations</th>
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<tr>
<td>Athletes with high-risk characteristics during EP study should undergo RF ablation to retain athletic eligibility.</td>
<td><img src="https://via.placeholder.com/15" alt="Heart" /></td>
<td>52, 69</td>
</tr>
<tr>
<td>An athlete may return to athletic activities after 3 months of successful ablation procedure in case of no recurrence of arrhythmia.</td>
<td><img src="https://https://via.placeholder.com/15" alt="Heart" /></td>
<td>52, 69</td>
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<tr>
<th>Athletes with implanted cardiac devices</th>
<th>Recommendations</th>
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<tbody>
<tr>
<td>Among athletes with ICD, performing an exercise test to determine the athlete’s upper heart rate for tachycardia zone programming is recommended.</td>
<td><img src="https://via.placeholder.com/15" alt="Heart" /></td>
<td>15, 73</td>
</tr>
<tr>
<td>Among the athletes with permanent pacemaker or ICD, only low–moderate intensity athletic activities except those with risk of bodily collision are recommended.</td>
<td><img src="https://via.placeholder.com/15" alt="Heart" /></td>
<td>15, 74</td>
</tr>
<tr>
<td>In asymptomatic athletes with Mobitz type 2 or complete AV block without structural heart disease, a deconditioning period up to 2 months is recommended. Persisting or recurring of symptoms after deconditioning may indicate pacemaker implantation.</td>
<td><img src="https://via.placeholder.com/15" alt="Heart" /></td>
<td>15, 73</td>
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<tr>
<td>Asymptomatic athletes with sinus bradycardia or sinus pauses that are secondary to elevated parasympathetic tone, permanent pacing should not be performed.</td>
<td><img src="https://via.placeholder.com/15" alt="Heart" /></td>
<td>15, 74</td>
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Presence of symptoms, ejection fraction, and arrhythmic status are the main determinants of an athlete with DCM who desires to participate in athletic activities.

Myocarditis
Myocarditis usually presents with symptoms of exertional dyspnea, chest pain, and arrhythmia. It easily mimics acute coronary syndrome, and coronary angiography is needed for the definite diagnosis (23). Autopsy studies in the series of sudden deaths in athletes indicate that myocarditis is a significant cause. Among the US military recruits, myocarditis-associated sudden death was found to be the most significant etiology (24).

Acute myocarditis can lead to DCM and can be resolved after a period of time with myocardial scar formation. This condition can lead to arrhythmias both during the acute and chronic phases. All physical activities should be restricted during acute myocarditis and exercise recommendations should be tailored after acute phase according to the clinical condition, laboratory, and imaging parameters of an athlete.

Arrhythmogenic right ventricular cardiomyopathy
ARVC is a hereditary myocardial disease caused by mutations, especially in genes encoding desmosomic proteins. ARVC is histologically characterized by the loss of myocytes in the right ventricular myocardium and/or LV myocardium with fibrofatty replacement which results in segmental or diffuse wall thinning (25).

The ECG findings of ARVC include most commonly T wave inversion in precordial leads V1 through V3 with a rare finding of Epsilon waves. The left bundle branch patterned ventricular tachyarrhythmia can be observed in ARVC patients (25).

Echocardiography and cardiac MRI may show right ventricular dilation or segmental wall motion abnormalities with morphological alterations. Fatty deposition in the right ventricular wall can be identified with cardiac MRI.

Both ventricular cavities can dilate physiologically in response to chronic exercise and this situation must be differentiated from the pathological conditions of ARVC. Generally, no segmental wall motion defects are seen in the physiological state. A positive family history along with ECG and echocardiographic findings may help identify ARVC, and typical findings on cardiac MRI confirms the diagnosis (26).

The most important risk factors in athletes with ARVC include prior history of SCD, sustained VT, or syncope (27).

Most of the SCDs in ARVC patients occur during exercise. Exercise itself can both facilitate the natural progression of the disease and lead to lethal arrhythmias. Reducing the intensity of the exercise in ARVC patients is associated with lower risk. Since exercise and SCDs have a definite causative relationship in ARVC, competitive athletic activities should be restricted in these patients.

Commotio cordis
Commotio cordis is defined as SCD due to ventricular fibrillation (VF) triggered by a blunt, non-penetrating blow to the precordial. Although initially considered extremely rare, it is now accepted as one of the most common causes of SCD in young athletes. Commotio cordis predominantly affects young male athletes (28).

Blows occur on the left chest wall and usually involve impact from a hard spherical object, such as a baseball, hockey ball, football, or volleyball. Strong body impacts which can occur during sports like karate and boxing may also lead to commotio cordis. Using soft balls or chest protection instruments during athletic activities can be useful but not absolutely protective (29).

More recently, the Commotio Cordis registry suggested that survival rates have increased steadily over the past 15 years, at >50%. Survival can be improved by earlier recognition of commotio cordis, shortening of the time interval from collapse to cardiopulmonary resuscitation, increased use of automated external defibrillators in the community, and increased number of people receiving training in cardiopulmonary resuscitation (30).

Coronary artery anomalies
The frequency of Coronary Artery Anomalies (CAAs) in the community is mostly learned by autopsies, angiography, and other imaging techniques (31, 32). Even though different results were also found in a number of angiography series, prevalence of CAAs have been reported between 0.21% and 5.79% in older surviving subjects (33, 34). Autopsy evaluation among military recruits and young athletes who experienced sudden death shows that CAAs are responsible for 17%–44% of cardiovascular mortality (31, 35).

In order of frequency among the general population, CAAs with an abnormal origin might present as the left circumflex artery originating from the right sinus valsala, a single coronary artery originating from the left sinus valsala, all coronary arteries originating from the right coronary sinuses, or the left anterior descending coronary artery (LAD) originating from the right coronary sinuses (36). Among athletes who have died suddenly, anomalous origin of LMCA and LAD from the right sinus valsala is more prevalent. The main clinical problem is the interarterial course of the left main coronary artery, LAD, and right coronary artery. The course of the vessels between the aorta and the pulmonary artery may cause compression of vessels, which in turn could lead to myocardial ischemia, ischemic arrhythmias, and sudden death (37-39).

Anomalous origin of left coronary artery from the pulmonary artery (ALCAPA) is usually an isolated anomaly. ALCAPA constitutes 0.22%–0.4% of congenital cardiac anomalies. Most patients die within the first year after delivery, so it is rarely seen among the athletes who have died suddenly (40).

The importance of preparticipation screening, especially in competitive young athletes, is evident in order to reduce mortality and morbidity caused by CAAs. Unfortunately, in contrast to congenital arrhythmia syndromes and cardiomyopathies, the diagnostic power of ECG in asymptomatic CAAs is not very high.

Echocardiography is superior to detect the concomitant structural congenital heart disease in athletes with CAAs. An experi-
enced echocardiographer can inform us about the origin of coronary ostia with 98% certainty. Transesophageal echocardiography can be performed if the images are not clear enough (41, 42).

Since our aim is to protect a young person from the negative consequences of sport activity, a maximal stress test must be performed in the third line after routine ECG and echocardiography in athlete with a possible diagnosis of CAA. If the test result is negative, it will carry him to a better level in terms of risk. If the stress test is suspicious, myocardial perfusion scan or stress echocardiography will be the best method to evaluate ischemia (43).

Recently, Multi Detector Computed Tomography (MDCT) data became recognized as an appropriate method to obtain the closest factual results. However, it is not practical to use MDCT as a routine control test due to restrictions related to the usage of X-ray and contrast agent (44, 45). Coronary angiography is indicated for definite diagnosis, and intracoronary functional assessment may be required.

The only treatment method for CAAs in athletes is surgery. Absence of myocardial ischemia must be demonstrated before returning to athletic activities in operated athletes.

### Channelopathies

#### Long QT syndrome

Cardiac ion-channel disorders, also known as “Channelopathies”, are inherited primary electrical disorders without “gross” cardiac structural abnormalities. Long QT syndrome (LQTS) is the most frequent within this group of suggestively pure “electrical” diseases, with an approximate prevalence of 1:2000 (46, 47). The disease is characterized by a prolongation of the QT interval (heart rate corrected QT) due to mutations in genes encoding for subunits of potassium, sodium, or calcium voltage-dependent ion-channels in the absence of secondary causes (48, 49). It is associated with syncope or SCD due to lethal ventricular arrhythmias (VAs), especially Torsades de Pointes, mainly triggered by adrenergic activation. The conditions associated with arrhythmic events are mostly gene-specific, with most arrhythmic events occurring during physical or emotional stress in LQT1, at rest or in association with sudden noises in LQT2 patients, and at rest or during sleeping in LQT3 patients (46, 47).

Restriction from virtually all competitive sports has formerly been the guideline-based recommendation since 2005 for athletes with any cardiac channelopathy (50, 51). However, starting from the “EHRA Expert Consensus Statement on the Inherited Primary Arrhythmia Syndromes” by Priori et al. (47) and 2015 AHA statement paper (52), expert panels tended to publish more liberal yet evidence-based recommendations (46, 47). ICD implantation is recommended in patients with previous SCD and in patients with syncope and/or for sustained VT occurring while receiving β blockers (46, 48).

In their milestone paper, Johnson et al. (53) sought for the outcomes of the global conventional exercise restriction rules suggested in LQTS. 70 athletes (54%) were competing contrary to European guidelines but within Bethesda guidelines (51, 54). However, none had a sport-related event. Of the 60 LQTS athletes (46%) continuing in sports contrary to both guidelines, only 1 experienced sporting-related events being equal to “1” event in “331 athlete-years”. In a large Italian registry young athletes with LQTS (0.6% of all non-eligible) were disqualified according to the contemporary guideline. During follow-up, no cardiac events in the disqualified athletes were reported. After more than 30 years of screening, the authors observed that only two of the LQTS cases had died suddenly (55). These results revealed that incidence of serious cardiac events in athletes with LQTS is lower than expected.

Athletes with LQTS should be supervised closely, and precautionary measures including treatment options should be implemented effectively.

#### Short QT syndrome

Short QT syndrome (SQTS) is a rare channelopathy characterized by a reduced duration of cardiac repolarization building the substrate for the development of lethal arrhythmias (46, 48). Five genes have been linked to SQTS (KCNH2, KCNQ1, KCNJ2, CACNA1C and CACNB2b), but the yield of genetic screening remains low (about 20% overall) (56). Resuscitated-SCD might be the first manifestation of the disease with a peak incidence in the first year of life (49, 56). SCD-survivors have a high recurrence rate; therefore, implantation of ICD is strongly recommended in this group of patients with/without quinidine or sotalol (46, 48). SQTS is diagnosed in the presence of a QTc ≤340 ms or QTc ≤360 ms and one or more of the clinical disease features (46). Limited data are available to quantify arrhythmic risk during competitive physical activity as well as genotype-phenotype relations in SQTS patients while even “syncope” seems to fail in predicting future events (46, 47). Individuals with SQTS should avoid dehydration, protein-supplements, excessive sweating, and hyperthermia during exercise.

#### Brugada syndrome

Brugada syndrome (BrS) is characterized by SCD and/or syncopal events due to VT/VF in young and apparently healthy individuals without significant medical history and with classical ST-segment-elevation-patterns in right precordial ECG leads (46, 47). BrS is inherited as an autosomal-dominant trait, which is more frequent in young adults and in men (57). The prevalence ranges from 1/1000 to 1/10 000 (46, 58). Either a decrease in the inward-sodium or calcium current or an increase in the outward-potassium-currents has been shown to be associated with the BrS phenotype. Ventricular-arrhythmia/SCD occurs at a mean 41±15 years, but it usually gets manifest during rest or sleep. ICD implantation is the definitive therapy in BrS patients with aborted SCD or with a history of cardiac-syncope and spontaneous type-1-pattern. The Brugada-pattern should carefully be distinguished from Brugada-phenocopies, which is challenging in an athlete’s ECG (59).

Data about the probable relation of exercise physiology and BrS is limited and mostly mechanistic rather than prognostic (60, 61).
In a meta-analysis (62) which included anecdotal cases; ST augmentation was observed during the early-recovery-phase of exercise in 57% of patients. There are insufficient data on the risks of exercise in BrS to make a recommendation. According to observations which suggest exercise might worsen the ST abnormalities in BrS and produce VA, patients with BrS might be restricted from vigorous exercise (60, 61). There is a risk of activation of temperature-dependent mutations at the climax of the exercise and sympathetic withdrawal in BrS patients. However, SCD in BrS occurs most often during sleep (60, 61). Altogether, this limited evidence might imply an already high-risk subgroup of BrS individuals manifesting their own poor-prognostic features instead of the detrimental effects of the “exercise physiology” itself.

Randomized and prospective data is certainly needed in order to reveal mechanistic relations and provide firm recommendations on cessation of sport participation. Until then, we lack crude evidence to expel all BrS population from exercising, apart from the high-risk-subgroup.

**Catecholaminergic polymorphic VT**

Catecholaminergic polymorphic VT (CPVT) is a rare, potentially life-threatening inherited arrhythmia with an estimated prevalence of 1:10,000 (63). It is diagnosed in the presence of a structurally normal heart, normal ECG, and unexplained exercise- or catecholamine-induced bidirectional VT or polymorphic ventricular premature beats or VT in an individual <40 years and/or in carriers of a pathogenic mutation in a CPVT-associated gene (47). Patients with CPVT often present with symptoms during the first decade of their life (63). Abnormal storage and release of calcium from the sarcoplasmic reticulum are the suggested mechanisms (64).

Intense physical activity has been implicated as a trigger for life-threatening cardiac arrhythmias in patients with CPVT. Exercise at a very low level should be allowed after the approval of CPVT experts (46). Appropriate precautionary measures should be taken during the physical activity of an athlete with CPVT including initiation of beta blocker therapy, electrolyte/liquid replacement, avoidance of dehydration, acquisition of a personal automatic external defibrillator, and establishment of an emergency action plan with the appropriate school or team officials (52).

**Early repolarization syndrome**

Early repolarization (ER) is a common ECG finding characterized by J-point elevation ≥1 mm ≥2 contiguous leads (47). The ER pattern in the precordial leads has been considered a benign phenomenon, but its presence in the inferior and/or lateral leads has been associated with idiopathic VF and/or polymorphic VT (ER syndrome) (47, 65). Non-anterior ER pattern including the inferior subtype is commonly seen in young competitive athletes (ranging from 14% to 44%) (66, 67). Noseworthy et al. (66) showed that both ER and the inferior subtype increased in prevalence with intense physical training. These data suggest that the ER pattern is a direct result of exercise training. ST-segment morphology variants associated with ER may help separate subjects with and without an increased risk of arrhythmic death in middle-aged subjects. Rapidly ascending ST segments after the J-point, which is the dominant ST pattern in healthy athletes, are not associated with an increased risk for arrhythmic death (68).

In conclusion, there is no evidence of an increased risk of SCD in healthy athletes with an ER pattern (63, 67, 68). Competitive sports may be allowed for a previously symptomatic athlete with ER syndrome with appropriate precautionary measures (52).

**Wolff Parkinson White syndrome**

The prevalence of Wolff Parkinson White (WPW) pattern is estimated to be 1–3 in 1000 individuals (69). Approximately 65% of adolescents and 40% of individuals over 30 years with a WPW pattern are estimated to be asymptomatic. Ventricular preexcitation accounts for approximately 1% of SCD in athletes (4). SCD may occur due to the development of atrial fibrillation (AF) with a rapid ventricular response that degenerates to VF (70). The main risk factor for SCD is the presence of an accessory pathway (AP) with short antegrade refractoriness (7).

Radiofrequency (RF) ablation should be performed in patients with WPW syndrome resuscitated from aborted cardiac arrest due to AF and rapid conduction over the AP causing VF (46, 69). If an athlete is symptomatic with syncope or palpitation, an electrophysiologic (EP) study is recommended. RF ablation is recommended if the refractory period of the AP pathway is ≤240 ms or the shortest preexcited RR interval is <220 ms during induced AF (46, 69). After 3 months of successfull ablation procedure, an asymptomatic athlete may return to athletic activities.

Approximately 65% of adolescents and 40% of individuals over 30 years with a WPW pattern are estimated to be asymptomatic (69). According to the 36th Bethesda Conference, an EP study is recommended for asymptomatic athletes if they participate in moderate- or high-level competitive sports (54). The ESC mandates that all athletes with WPW undergo an EP study for risk assessment (52). Asymptomatic patients with a short preexcited RR interval ≤250 ms in AF or <220 ms during stress or isoproterenol, the refractory period of the AP ≤240 ms, presence of multiple APs, or easily induced AF are at increased risk for SCD (47, 69). Athletes with high-risk characteristics mentioned above during EP study should undergo RF ablation to retain athletic eligibility. In addition, asymptomatic patients with WPW and structural heart disease or ventricular dysfunction secondary to dyssynchronous contractions may be considered for ablation regardless of the antegrade characteristics of the AP (69).

**Athletes with implanted cardiac devices**

**Athletes with permanent cardiac pacemakers**

Highly-trained endurance athletes have dominant parasympathetic tone at rest associated with marked sinus bradycardia,
first degree and Mobitz type I atrioventricular block. Mostly these findings are physiological events that do not require intervention (71). Underlying structural heart disease should be excluded in case of Mobitz type 2 atrioventricular block or third-degree atrioventricular block (72).

Potential risk for device and lead damage may limit the professional life of athletes. The 36th Bethesda Recommendations state that PM dependent athletes should not participate in sports that can involve bodily trauma (1). Although not addressed in formal recommendations, protection with padding might be considered. Programing upper tracking rates at higher levels is important in athletes with complete heart block. Heart rates of the patient during vigorous exercise should be considered. Myopotential inhibition may lead to inhibition of pacing, which is of concern in pacemaker-dependent patients. Therefore, bipolar leads should be selected in athletes.

Athletes with ICD

It should be kept in mind that ICDs do not prevent the occurrence of ventricular arrhythmias and do not effect the progression of the underlying disease. Therefore, careful consideration of the underlying disease is mandatory before participation in competitive sports (15).

However, a long-term prospective multinational registry provides promising data about this topic. After a mean follow-up period of 44 months of 440 patients, there were no arrhythmic deaths, externally resuscitated tachyarrhythmias during sports participation, or injury resulting from arrhythmia-related syncope or shock during sports (73). 31 definite and 13 possible lead malfunctions were reported. The estimated lead survival free of definite plus possible malfunction was 94% at 5 years and 85% at 10 years. No generator malfunction was reported.

On the other hand, approximately one in five received both appropriate and inappropriate shocks, which mainly occurred during competition or physical activity. Therefore, programing an ICD of an athlete is always challenging due to the risk of inappropriate shocks from high heart rates during exercise. Recently, the role of ICD programming characteristics on occurrence of shocks, transient loss of consciousness, and death among athletes was assessed by prospective, observational, international registry (74). High-rate cutoff and long-detection duration programing in athletes was associated with reduction in total and inappropriate ICD shocks without affecting survival or the incidence of transient loss of consciousness. Since it is not easy to recommend certain tachycardia detection zones, patient-tailored programming seems a better approach.

The potential risks associated with mechanical trauma are possible for ICDs. The athlete’s ability to participate in sports should be discussed individually. Underlying cardiovascular disease, type and the programing of the device, type of the sport, risk for trauma, and risks related to potential syncope or shock should be considered.

Conclusion

Specific return-to-play protocols should be developed for competitive athletes following treatment of various cardiovascular conditions like catheter ablation, cardiac device implantation, and corrective surgeries.

Athletic activities performed in a competitive fashion could have hazardous effects on the cardiovascular health of the athletes. Athletes who have high-risk genetic cardiovascular diseases and implanted cardiac devices should be closely supervised by sports cardiologists, exercise specialists, and their personal trainers in a deep collaboration with the guidance of professional and scientific recommendations.

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