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Myocarditis in athletes is a challenge – diagnosis, risk stratification and uncertainties

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Abstract

Presentation of myocarditis in athletes is heterogeneous and establishing the diagnosis is challenging with no current uniform clinical gold-standard. The combined information from symptoms, electrocardiography, laboratory testing, echocardiography, cardiac magnetic resonance imaging (CMR), and in certain cases endomyocardial biopsy (EMB) helps to establish the diagnosis. Most patients with myocarditis recover spontaneously; however, athletes may be at higher risk of adverse cardiac events. Based on scarce evidence and mainly autopsy studies and expert’s opinions, current recommendations generally advise abstinence from competitive sports ranging from 3 to 6 months. However, the dilemma poses that (un-)necessary prolonged disqualification of athletes in order to avoid adverse cardiac events, can cause considerable disruption to training schedules and tournament preparation, and leading to a decline in performance and ability to compete. Therefore, better risk stratification tools are needed. Using latest available data, this review contrasts existing recommendations and presents a new proposed diagnostic flowchart putting a greater focus on the use of CMR imaging in athletes with suspected myocarditis. This may enable cardiac caregivers to risk stratify athletes with suspected myocarditis more systematically, and furthermore allow for pooling of more unified data. To modify recommendations regarding sports behavior in athletes with myocarditis, more evidence, based on large multicenter registries including CMR and EMB, is needed. In the future, physicians might rely on combined novel risk stratification methods, by implementing both noninvasive- and invasive tissue characterization methods.

Key words:
Myocarditis; Infection; Athletes; Sports; Recreational; Competitive, Sports Restriction; Cardiac Magnetic Resonance Imaging; Risk Stratification

Abbreviations:
BNP = Brain Natriuretic Peptide
CAD = Coronary Artery Disease
CMR = Cardiac Magnetic Resonance Imaging
ECG = Electrocardiography
ECV = Extracellular Volume Fraction
EMB = Endomyocardial Biopsy
LGE = Late Gadolinium Enhancement
LLC = Lake Louise Criteria
LVEF = Left Ventricular Ejection Fraction
MACE = Major Adverse Cardiovascular Events
RV = Right Ventricular
RVEF = Right Ventricular Ejection Fraction
SCD = Sudden Cardiac Death
TTE = Transthoracic Echocardiography
Introduction

Myocarditis is an underlying cause of sudden cardiac death (SCD) in young athletes (1-3). While the beneficial effects of exercise on both cardiovascular and general health are irrefutable, some evidence has emerged showing that excessive levels of exercise in the presence of an infection and particularly in myocarditis can be harmful (2,4). Current recommendations state that after a case of myocarditis abstinence from competitive sports lasting between 3 to 6 months is generally recommended; this, however, can be extended to up to 1 year and is based on scarce evidence, mainly autopsy studies and experts’ opinions (5-7). However, prolonged disqualification of athletes is disruptive to athletic conditioning, which may lead to performance drop and inability to compete. In order to avoid adverse cardiac events and better risk stratify athletes with myocarditis, prognostic information may be gained from the application of using cardiac magnetic resonance imaging (CMR) (8). This review presents a new proposed diagnostic flowchart (Central Illustration) based on recent evidence and existing recommendations, which integrates the use of CMR imaging for diagnosing and risk stratification of athletes with suspected myocarditis.

Myocarditis in Athletes

In post-mortem studies of athletes who died from SCD, myocarditis was diagnosed in up to 8% (9,10). This represents the third most common cause after autopsy-negative sudden unexplained death and coronary artery anomalies (1). It is uncertain if physical activity is truly necessary to trigger malignant arrhythmias as the majority of SCD, especially due to myocarditis, did not occur during or immediately after exercise but at rest (1,11). Viral infections, namely with enterovirus, Coxsackie B, Parvovirus B19 and Human Herpes Virus 6 are the most common responsible infectious pathogens in myocarditis (12,13). Initially,
these pathogens directly affect the myocardium, followed by autoimmune myocardial injury, which ultimately results in myocardial remodeling potentially leading to dilated cardiomyopathy. Another factor that determines the underlying causative pathogen may be governed by the type of sports performed. For example, cross-country runners have greater exposure to tick-bite-transmitted infections (Borrelia Burgdorferi). Myocarditis can also be caused by a great variety of medical drugs as well vasculitic or toxic causes (14), underlining the attention of considering co-morbidities in athletes. In general, non-infectious causes of myocarditis are rare in athletes but can include illicit drugs (15), illustrating the importance of lifestyle history in athletes.

Pathogenesis and Contributing Effect of Strenuous Exercise

The general hypothesis is that moderate exercise, which does not comprise of a common definition amongst studies of runners, is protective. However, more intense endurance training, typically that for a marathon may increase the likelihood of a systemic inflammatory response and susceptibility to viral upper respiratory tract infections (16). As a result, acute myocarditis can ensue and lead to an interaction between an environmental trigger and the host’s immune system, leading to three phases of disease progression: acute viral, subacute immune and chronic phase, with 12 - 25% developing dilated cardiomyopathy (17). While the viral phase is often short-lived and not clinically detected, pathogens may reach the myocardium and affect it through direct myocyte injury as well as through activation of the innate immune system; this results in subacute and chronic inflammation, which leads to myocyte necrosis, fibrosis and ultimately adverse cardiac remodeling (17), pre-disposing an individual to arrhythmias. Exercise, particularly under extreme physical exertion, is associated with a higher propensity to arrhythmogenicity (18), which murine studies have shown to be due to an intensified inflammatory response in subjects exposed to extreme physical exertion (18,19). Even after the
myocardium has recovered after an episode of myocarditis, the chronic myocardial inflammatory process secondary to pathogenic autoimmunity may persist, where cytokines carry on to exert pro-arrhythmic effects and trigger circuit activity (20). Therefore, athletes with a history of myocarditis, or in certain cases even just a minor infection, may pose an increased risk for adverse cardiac events if physical exercise is continued.

**Clinical Presentations and the Use of Biomarkers, Viral Serology, Electrocardiogram and Transthoracic Echocardiography**

Myocarditis presents a challenging clinical scenario given its heterogeneous presentation. The suspicion of myocarditis may be raised when athletes complain of chest pain or indeed any other cardiac symptom in the context of general malaise with abnormal biomarkers, ECG or transthoracic echocardiogram (TTE) changes. This particularly affects those individuals who present with new onset left ventricular (LV) dysfunction after viral infection or prodromal symptoms in recent weeks. One must be aware that signs of myocarditis may resemble those of physiological changes in the athlete’s heart (Table 1).

**Biomarker and Viral Serology**

Other than an increase in troponin, there are no other specific biomarkers indicative of myocardial injury due to myocarditis. Evidence for viral serology has not proved sufficient as shown by a prospective study by Mahfoud et al., which determined the sensitivity and specificity of virus serology to be 9% and 77% respectively. Only 5 out of 124 patients were shown to have viral serology consistent with the virus that was detected upon EMB (21). Where there is clinical suspicion or high risk, testing for hepatitis C, human immunodeficiency virus, Lyme disease or rickettsia is warranted. A recent study characterizing the causes of raised troponin in those 50 years or less including 6081 patients showed that myocarditis was the second most common
cause of raised troponins after myocardial infarction (22), further underlining the importance of myocarditis as a diagnosis especially in young and middle-aged patients. A meta-analysis including thirty-three studies with a total of 1045 athletes, mainly participating in endurance sports ranging from short runs to ultra-marathons showed that cardiac troponin was elevated above the 99th percentile in up to 83% of individuals following prolonged exercise (23). However, the troponin release tends to be less prominent and monophasic rather than biphasic and resolves faster than in myocardial injury as caused by myocarditis or acute coronary syndrome. Therefore, in the absence of an acute coronary syndrome, a judgement on the nature of the myocardial insult should consider the prominence of the rise in troponin, the timeframe of a high troponin level and the time elapsed since endurance exercise. Nevertheless, a recent study by Berg et al. has concluded that absolute baseline levels of cardiac enzymes and inflammatory biomarkers do not sufficiently represent the level of Late-Gadolinium Enhancement (LGE) by CMR in myocarditis. Further, it does not predict change in LGE at 3 months follow-up and does not correlate with improved LGE when a decrease in cardiac enzymes and inflammatory biomarkers is observed (24). Based on their findings, they concluded that it was not sufficient to simply use clinical findings, cardiac enzymes and inflammatory markers to monitor myocarditis and that CMR adds important additional information to current diagnostic techniques.

Electrocardiogram

Patients with myocarditis may present with unspecific ST-elevations, PQ-depression, QTc prolongation or T-wave inversion, which, in some cases, might be difficult to distinguish from normal variants in healthy athletes (Table 1) (25,26). In our recent report looking at the prognostic value of CMR tissue characterization in risk stratifying patients with suspected
myocarditis, only 42% of the 670 suspected myocarditis patients showed abnormal ECGs, with no significant difference between those where LGE was present and absent (8).

**Transthoracic Echocardiography**

Generally, the heart of trained athletes can present with various structural changes depending on the dynamic component of the type of exercise performed (27) – endurance training, such as that for a marathon, increases isotonic load on working muscles that can lead to increased LV dimensions, while high isometric load on working muscles, such as in weight lifting, can result in increased LV wall thickness. To what degree different types of exercise involve dynamic and static components has been further described by Mitchell et al. (28) and may help physicians to counsel athletes with cardiovascular abnormalities in regards to particular types of competitive exercise. Compared to healthy athletes, in a case of myocarditis, there are no specific features in TTE. Myocarditis can resemble dilated, hypertrophic, or restrictive cardiomyopathy with local wall motion abnormalities and/or pericardial effusion (29).

Nonetheless, TTE can be of prognostic value in fulminant myocarditis, which presents with normal chamber size and severe impaired systolic left ventricular ejection fraction (LVEF) (30). It has been shown that regional wall motion abnormalities following myocarditis in athletes can be exacerbated by exercise in contrast to normal-/depressed LVEF among healthy athletes, which normalizes during exercise (31). TTE is therefore an important and feasible tool to fully assess LV function and wall motions at presentation and at follow-up. The LV remodeling index was identified to differentiate athletes from patients presenting with dilated and hypertrophic cardiomyopathy (32). However, TTE cannot always help to differentiate athlete’s heart from other pathological structural or functional changes and in those cases CMR may be indicated as it also helps to identify further diagnostic information beyond geometric measures (33).
Invasive Coronary Angiography and Endomyocardial Biopsy

Generally, coronary artery disease (CAD) should be ruled out first using invasive or noninvasive imaging or a clinically low-pretest probability for CAD in individuals presenting with suspected myocarditis. If at the time of CMR scanning, no signs of active inflammation on T2-weighted imaging or T2 mapping is present, clinicians could consider performing CMR stress-perfusion to rule out ischemia. In this setting, CMR may further help to differentiate between an ischemic event with its typical pattern of endocardial LGE presence in coronary vessel distribution territory versus epicardial LGE in myocarditis (e.g. in athletes with low pre-test probability and no indication for invasive coronary angiography). Concerning the use of endomyocardial biopsy (EMB), European and American recommendations differ slightly (Table 2). EMB is an invasive diagnostic test with high specificity and complication rates as low as <1% in experienced centers (34,35). EMB can differentiate between different types of inflammation (infectious, autoimmune, idiopathic) causative of myocarditis and may therefore guide treatment and prognosis. This is particularly recommended in life-threatening presentations or unexplained reduced systolic LV function (14). However, it lacks sensitivity and exhibits high false negative results (sampling error). To avoid focal sampling error, EMB is preferably performed soon after presentation and multiple samples of sufficient size (1 – 2 mm) should be taken from both ventricles (35,36). Additionally, diagnosis by EMB can be improved by analyzing the viral genome through DNA-RNA extraction and RT-PCR amplification (37). This has the advantage of knowing exactly which pathogen is causing disease as different viruses have different effects on myocardial and vascular tissue and affect different areas of the myocardium, which in turn may affect presenting signs and symptoms as well as ECG. Historically, it has not been clear whether CMR helps in guiding biopsy and improving
sensitivity/specificity of EMB (35,38). A study by Baccouche et al. suggests a diagnostic
synergy in using both CMR and EMB as complementary diagnostic tools in troponin-positive
patients without CAD, identifying 78/82 myocarditis patients (95%) when applied together,
which was superior to both CMR and EMB when applied individually (39). Recent evidence of
real-time CMR-guided EMB in a porcine pre-clinical in-vivo model suggest that CMR-guidance
of EMB may significantly improve sensitivity and specificity of EMB (40). However, whether
EMB would help to earlier risk stratify athletes with regard to sports abstinence counseling is
unknown. Therefore, performing EMB generally has the greatest potential benefit in cases where
CMR cannot confirm a diagnosis of myocarditis or in cases with unclear depressed LVEF
(Central Illustration).

**Cardiac Magnetic Resonance Imaging**

CMR plays a major role in evaluating the etiology of chest pain syndromes and suspected
myocarditis but recommendations about the exact role of CMR in diagnosis and follow-up of
myocarditis (Table 2) (41) are outdated due to new data and improvements in imaging technique.
CMR has evolved as the primary noninvasive diagnostic modality in suspected myocarditis
cases, with particularly high sensitivity (81%), specificity (71%) and diagnostic accuracy (79%)
in acute myocarditis (42), given several technical advantages. Beyond the assessment of wall
motion and LVEF, CMR is a noninvasive modality that allows tissue characterization with
visualization of myocardial edema and fibrosis (43).

The Lake Louise Criteria (LLC) (44) utilized in the diagnosis of suspected myocarditis
combine different CMR techniques to detect intramyocardial edema by T2-weighted imaging,
fibrosis by LGE, typically in a non-ischaemic distribution, as well as hyperemia by myocardial
early gadolinium enhancement (EGE). A certain number of patients with biopsy proven
myocarditis have no LGE and do not fulfil the LLC (35,45); reasons for this may include that active myocarditis may not always lead to regions of necrotic myocytes that are large enough to be detected by CMR. Moreover, the optimal time point after onset of symptoms for a CMR scan has not yet been determined yet and (too) early CMR scanning may yield false negative findings which necessitate repeat scanning (Central Illustration).

CMR may detect isolated myocarditis-like LGE on CMR in an athlete with non-acute symptoms (e.g. syncope, or palpitations without any clinical signs of acute myocarditis). Whether these findings represent an acute myocarditis, a myocarditis in a late stage without presence of myocardial edema, a resolved myocarditis or another pathology can be challenging (see Figure 2), however, CMR can be considered the most suitable noninvasive modality for differentiation of different underlying cardiac causes.

T2 weighted imaging, T1 - and T2 mapping techniques

LLC are now often complemented by novel CMR techniques such as native T1 and T2 mapping as well as extracellular volume fraction (ECV) measurement to provide a higher sensitivity and specificity (46,47). Native T1 mapping values are affected by both edema and extracellular expansion and is therefore able to detect myocarditis at various stages, whereas T2 mapping evaluates free water content that is normally present in the acute phase of myocarditis and then gradually normalizes over months - this is relevant for cardiac caregivers evaluating the course of myocarditis as it is the only technique that can adequately discriminate between myocarditis and noninflammatory cardiomyopathies in patients with symptoms lasting longer than 2 weeks (48,49). A recent meta-analysis by Pan et al. (48) comparing the diagnostic performance of ECV, native T1 mapping and T2 mapping versus LLC for detection of acute myocarditis including 17 studies and 1308 subjects, has shown that only native T1 mapping had
significantly better sensitivity than LLC, while other parameters had comparable diagnostic performance to LLC and provided distinct advantages for evaluating myocarditis. The meta-analysis shows that LLC have been firmly established in clinical practice and that the use of native of T1 mapping can further enhance CMR accuracy in diagnosing myocarditis. These findings are supported by another meta-analysis including 22 studies by Kotanidis et al. (50), which similarly concludes that native T1 mapping has a significantly higher diagnostic accuracy compared to all other index tests: native T1 mapping showed an area under the curve (AUC) of 0.95 compared to T2 mapping with an AUC of 0.88 and LLC of 0.81 and will therefore likely be implemented in the upcoming updated Lake Louise Criteria. Additionally, the authors showed that T2 mapping was superior to T2-weighted and EGE imaging, suggesting that both EGE and T2-weighted imaging could be replaced by T1 - and T2 mapping respectively (50). The role of ECV is less clear at this stage. In our subset analysis, ECV showed a significant association with major adverse cardiovascular events (MACE) and death and may therefore be of additional benefit in a routine CMR for risk stratification in suspected myocarditis (8).

However, due to its novelty, it is important that further studies on the inherent characteristics of T1 mapping will be performed. As T1- and T2 mapping values are vendor and site specific, standardization of these mapping techniques and understanding how confounding factors will influence them in real-life clinical practice outside of the research setting is needed (51).

Furthermore, native T1 prolongation becomes less specific to myocarditis over time as early inflammation is replaced by fibrotic tissue, which, in patients with chronic symptoms, can be due multiple cardiac pathologies. To truly implement changes to diagnostic criteria, head-to-head comparisons performed at multiple centres would be desirable (52).
In our own sub-group analysis, T2 weighted imaging has demonstrated a significant association with MACE (8). Other studies were not able to show this predictive value, possibly due to the low patient number and the fact that T2 weighted imaging is prone to artefacts. Our study is supported by the study of Spieker et al. who could show that myocardial edema detected with T2 mapping has prognostic value in patients with myocarditis. Furthermore, it seems that T2 mapping is more robust and less prone to artefact compared to T2 weighted imaging and may be the better sequence for edema assessment in patients with suspected myocarditis (53,54).

Late Gadolinium Enhancement

Whether any LGE presence in athletes may be the substrate for ventricular tachycardia (e.g. from previous myocarditis or other causes) is unknown (55,56). Studies have reported a high rate of coincidentally detected LGE in athletes (55-57). In two recent series, which included athletes who had undergone CMR because of abnormal screening (ECG changes or ventricular arrhythmias) first line examination but structurally normal hearts on echocardiographic imaging, a high rate of cardiac events during follow-up was reported in those with large areas of sub-epicardial LGE (58,59). Although these athletes had no definite history of myocarditis, the authors hypothesise that due to the typical non-ischemic epicardial pattern of LGE a silent myocarditis is most probably the underlying cause. Although rather untypical, other causes like arrhythmogenic right ventricular cardiomyopathy (ARVC) with mainly left heart involvement are not completely excluded, however, it is generally more likely that myocarditis mimics an ARVC than vice versa (60). Furthermore, in the acute stage differential diagnosis of myocarditis and ARVC does not change the sports recommendation, but does change diagnostic downstream testing. The mentioned studies highlight the fact that subepicardial fibrosis may otherwise not have been detected by echocardiography alone and that CMR plays an important role especially
in athletes with ventricular arrhythmias. These findings support the argument to use CMR early for risk stratifying athletes with suspected myocarditis, which, when LGE is present, most commonly presents with subepicardial LGE (8,61). Our group had recently shown that in 670 patients with suspected myocarditis with a median follow-up of 4.7 years, CMR is not only a powerful diagnostic but also a prognostic tool (8). In fact, patients with suspected myocarditis and presence of LGE (n = 296, 44%) showed a doubled risk for MACE at follow-up. Regarding location and pattern, septal and mid-wall LGE demonstrated strongest significant associations with MACE (Hazard Ratio of 2.6 and 2.4). A patchy distribution portended to a near 3-fold increased hazards to MACE and LGE extent (increase by 10%) corresponded to a near eighty percent increase in risk of MACE. The Italian Multicenter study on Acute Myocarditis (ITAMY), which included 374 patients with acute myocarditis and preserved LVEF, similarly concluded that in these patients LGE in the mid-wall layer of the anteroseptal myocardial segment is associated with a worse prognosis (61). Mahrholdt et al. found that varying patterns of LGE are associated with different viruses; parvovirus B19 was associated with inferolateral LGE, while anteroseptal LGE was associated with either human herpesvirus 6 or the combined presence of the two viruses (62). On the contrary, a normal CMR study corresponded to low annual MACE and death rates of 0.8% and 0.3% respectively in our study (8), and no events were recorded in the ITAMY study (61). Alluding to the diagnostic usefulness and common application of CMR, our study showed that only 57 (9%) of patients underwent EMB with mostly unspecific results, which is consistent with other studies (44,63,64), while the ITAMY study (61) solely relied on LLC for diagnosis of acute myocarditis in 95% of cases. The prognostic power of LGE presence on CMR in suspected myocarditis is similarly shown by both our study (8) and the ITAMY study (61). It is to note that our study shows lower event rates as
we did not select patients who have already been diagnosed with acute myocarditis according to LLC; rather we chose a more real-life clinical setting, where low-risk patients are more likely to be included in the patient population; additionally, ITAMY and our study showed slight differences in MACE criteria.

Another important risk factor that will require further studies evaluating its prognostic strength is LV remodeling. Recently, Filippetti et al. (65) showed that not only the lack of LGE but also of any adverse LV remodeling at mid-term control scanning resulted in better outcomes in acute myocarditis.

CMR used as a diagnostic and risk stratification tool can help to identity how physicians should approach treatment of an athlete with current or recent infection and suspected myocarditis, what to recommend regarding disqualification from exercise and may help to fill apparent evidence gaps (Central Illustration). Assuming that cardiac centres have sufficient CMR experience and image quality is adequate, CMR can act as a gatekeeper in the diagnosis of myocarditis and a revised diagnostic algorithm that incorporates CMR in a uniform fashion will help to create necessary outcome data. While athletes with LGE presence may need to undergo CMR follow-up, absence of LGE on CMR scanning in athletes with suspected myocarditis could be interpreted as ‘low-risk’ and they might get allowed to go back to exercise after clinically resolved inflammation (66).

**Therapy**

In most cases, myocarditis is a benign pathology and resolves favorably. Currently, no randomized controlled trials for optimal therapy exist. Current recommendations (14) endorse heart failure medication as supportive therapy for LV dysfunction in myocarditis. Beta-blockade improves LVEF and reduces hospitalization while increasing survival; carvedilol has been
shown to exert cardioprotective effects (67), while metoprolol may play a role in increased inflammation (68). Non-steroidal anti-inflammatory agents are generally only advisable in patients with peri-/myocarditis with normal LVEF and chest pain (69). The use of immunosuppression with a combination of cortisol and azathioprine for 6 months has shown benefits in a single-centre study when there is no viral persistence (70). However, current recommendations do not include these immunomodulatory therapies. Of note, immunosuppression may be useful and improve outcome in patients with a particular form of myocarditis, a giant cell myocarditis (71). Beside heart failure therapy, supportive arrhythmia management is recommended. Specific recommendations for arrhythmias as well as intracardiac defibrillator implantation in myocarditis do not exist and hence, management should be in concordance with current arrhythmia guidelines. Wearable life-vests (72) may play a future role in athletes, who may want to try getting back into some low intensity exercise at an earlier stage, even at an increased risk of SCD since the effect of adverse remodeling will not be mitigated (if the physical aspects of the type of sports allow it, e.g. spinning).

**Follow-up and sports restriction**

First and foremost, athletes with acute myocarditis need to refrain from physical exercise, especially from competitive sports. Based on the most recent scientific statement from the AHA/ACC, it is recommended that athletes with probable or definite diagnosis of myocarditis should not participate in competitive sports while active inflammation is present (Class III, Level of evidence C) (Table 3), signs of which are most easily detected using CMR (e.g. T2-weighted imaging for edema). This recommendation applies across patient age groups, genders and left ventricular functions. Before returning to competitive sports, athletes should undergo an echocardiography, Holter monitoring – preferentially before and during a work-out, and an
exercise ECG no less than 3 - 6 months after initial illness (Class I, Level of Evidence C). There is currently no evidence available on the ‘safe’ level of exercise in the 3 - 6 months “competitive sports abstinence” period in athletes with myocarditis. High intensity training would be rather categorized as competitive sports and should not be recommended (see case presentation in Figure 1) during this period. Yet, the role of moderate or isometric exercise is still unclear. It is an individual case by case decision and according to the “Exercise in heart failure” consensus document (73) moderate exercise at 50% VO2peak or 60% from their maximum predicted heart rate is recommended and may also be translated to myocarditis patients. However, clinical and laboratory absence of inflammation and absence of arrhythmias are a requirement to return to any exercise levels. Other missing gaps include whether different levels of ‘severity’ of myocarditis or locations, patterns and size of LGE require different lengths of sports abstinence and whether serial CMR scans are needed to guide therapy and recommendation regarding sports behavior. Furthermore, to what extent genetic predisposition determines disease progression and how clinical examination, biomarkers, ECG and EMB can help to anticipate adverse outcomes is still unclear.

CMR has the added advantage to unveil possible acute myocarditis by identifying myocardial edema even in absence of LGE. In order to avoid adverse cardiac events, this may allow clinicians to make conservative recommendations regarding returning to exercise until inflammation is resolved of the myocardium in the follow-up CMR (see Central Illustration). In a small study evaluating 28 myocarditis patients (non-athletes), patients with decreasing LGE at follow-up had a lower event rate compared to those with greater amount of LGE (74). However, at present it remains unresolved if resolution of myocarditis-related LGE should be a requirement prior to returning to competitive sports (5).
Recommendations should be applied not only to competitive athletes but also to those participating in recreational sports. Two studies showed that most myocarditis related SCD most frequently occurs in recreational athletes (75,76). Current European recommendations reflect this in that they generally recommend no competitive sports for those diagnosed with myocarditis (6), while also extending this to recreational and amateur sports activities (7,14); clinical assessment in the absence of abnormal LVEF and arrhythmias is necessary prior to resuming competitive sports. CMR features such as T2-weighted imaging, LGE and T1/T2 mapping with regard to sports behavior recommendations are currently not included. Further research will be needed in order to assess how this novel mapping techniques can help treating physicians to counsel athletes with myocarditis.

**Conclusion**

Information leading to sports restriction recommendations in athletes with myocarditis are mainly based on autopsy studies, animal models and experts’ opinions. Recent evidence showed that CMR as a non-invasive imaging tool plays an important role in the risk stratification of patients with suspected myocarditis. To modify recommendations regarding sport behavior in physically active individuals with myocarditis, more evidence, based on large multicenter registries including CMR and immunochemistry EMB, is needed. Physicians might rely in the future on combined novel risk stratification methods, which will be likely improved by implementing both noninvasive- and invasive tissue characterization methods using CMR and EMB.
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### Tables

**Table 1. Challenges in ECG, biomarkers and noninvasive imaging characteristics in patients with athlete’s heart versus acute myocarditis**

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<thead>
<tr>
<th></th>
<th>Differentiating Features</th>
<th>Differentiating Features</th>
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<td></td>
<td>Athletes Heart</td>
<td>Myocarditis</td>
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<tr>
<td>Symptoms</td>
<td>Asymptomatic</td>
<td>Symptomatic</td>
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<tr>
<td>ECG/Holter</td>
<td>1. <strong>Specific ECG</strong> changes such as early repolarization/ST segment elevation, T-wave inversion in V1-V3 ≤age 16 years old, ST elevation followed by T wave inversion V1-V4 in black athletes</td>
<td>1. <strong>Unspecific ECG</strong> changes. Possible PQ depression, ST elevation in multiple leads.</td>
</tr>
<tr>
<td>Biomarkers/Inflammatory markers</td>
<td>1. <strong>Troponin elevation</strong> mild and normalizes quickly. May be present in ultra-endurance athletes.</td>
<td>1. <strong>Troponin elevation</strong> mild to high</td>
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<td></td>
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<td>2. <strong>Others</strong>: BNP elevation, Creatine-Kinase, Leucocytosis,</td>
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<td></td>
<td><strong>TTE</strong></td>
<td><strong>CMR</strong></td>
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<td><strong>2. Others:</strong> BNP mildly elevated after ultra-endurance exercise</td>
<td>elevated C-reactive Protein, elevated Erythrocyte sedimentation rate</td>
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<td><strong>1. Ejection fraction:</strong></td>
<td><strong>1. Ejection fraction:</strong></td>
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<td>Sub-depressed LVEF and/or RVEF in ultra-endurance athletes, normalizes</td>
<td>Depressed LVEF, can further decline when exercising.</td>
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<td>when exercising.</td>
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<td></td>
<td><strong>2. Dilatation and eccentric remodeling,</strong></td>
<td><strong>2. Focal hypokinesia</strong></td>
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<tr>
<td></td>
<td>no focal regional wall motion abnormalities</td>
<td>at rest or during exercise (regional wall motion abnormalities)</td>
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<td></td>
<td><strong>3. Pericardial Effusion</strong></td>
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<tr>
<td></td>
<td><strong>1. LGE typically absent;</strong></td>
<td><strong>1. LGE:</strong> normal or specific mid to sub-epicardial LGE pattern</td>
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<td>possible LGE if previous silent myocarditis</td>
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<td></td>
<td><strong>2. Edema rarely present</strong></td>
<td><strong>2. Edema present</strong></td>
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<td>after exercise</td>
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Table 2: Recommendations regarding endomyocardial biopsy and the use of CMR in patients with suspected myocarditis

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<td>Presents 13 scenarios in which EMB might be considered. Class IB recommendations for ‘EMB in the setting of unexplained, new-onset heart failure of… &lt;2 weeks’ duration associated with a normal-sized or dilated left ventricle in addition to hemodynamic compromise.’</td>
<td>Use EMB widely in order to make a diagnosis, management plan and prognosis according to whether viral genomes and inflammation are present.</td>
<td>Routine EMB in all cases of heart failure is not recommended. In those patients progressing rapidly or suffer from unexplained cardiomyopathy and where active myocarditis, in particular giant cell myocarditis is suspected, should undergo EMB.</td>
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And

… 2 weeks’ to 3 months’ duration associated with a dilated left ventricle and new
ventricular arrhythmias, Mobitz type II second- or third-degree atrioventricular (AV) heart block, or failure to respond to usual care within 1 to 2 weeks.’

**Differences:**

European and US guidelines differ in so far as European guidelines do recommend routine EMB in myocarditis cases, whereas US guidelines are more conservative in using EMB.

**Recommendations regarding the use of CMR in patients with suspected myocarditis**

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<td>Recommends the use of CMR as part of a more comprehensive diagnostic approach. Particularly in patients with significant ongoing, recurring or persisting inflammation, it is able to determine the extent and regional distribution of reversible and irreversible</td>
<td>Reasonable to perform CMR prior to EMB but only if the situation is not life-threatening. CMR should not replace EMB as a diagnostic tool. The use of the ‘Lake – Louise – Criteria’ is recommended.</td>
<td>Pericardial effusion on CMR or characteristic alterations in tissue signal on T2- or T1-weighted images and the presence of LGE are sufficient for the diagnosis of probable acute myocarditis if a clinical syndrome that includes acute heart failure, angina-type chest pain, or</td>
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</table>
myocardial injury, as well as to detect functional abnormalities. Proposes ‘Lake – Louise – Criteria’. Pooled diagnostic accuracy of 78% with a sensitivity of 67% and specificity of 91% for diagnosis of myocarditis on CMR compared to biopsy as the gold-standard.

| Differences: |
| US recommendations are more inclusive of CMR as a diagnostic tool and do not require EMB in the assessment of myocarditis, whereas European recommendations include the use of CMR only complimentary to EMB. |

myopericarditis of <3 months’ duration is present.
Table 3: Sports Restriction Recommendations in athletes with myocarditis

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<td>Medical history, physical</td>
<td>Biopsy not required to guide clinical management but recommended in certain cases (see Table 2). Time period of absence from exercise will depend on severity of initial symptoms. Based on experimental models, a resting period of at least 3 – 6 months is recommended. No exercise during active inflammation for patients with probable or definite myocarditis. Within 3 – 6 months of presentation, ECG, 24-hr Holter and exercise ECG to assess fitness to exercise – normal systolic function, normal biomarkers and absence of arrhythmias.</td>
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<td>examination, 12-lead ECG and</td>
<td>Temporary exclusion of both competitive and recreational athletes independent of age, gender and severity for a minimum of 6 months. Clinical assessment prior to resumption required.</td>
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<tr>
<td>echocardiography.</td>
<td>Medical history, physical examination, 12-lead ECG and echocardiography according to individual case assessment.</td>
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<td>No competitive exercise during</td>
<td>No competitive exercise during active myocarditis or pericarditis. First control within 6 months through assessment by the above modalities as well as exercise testing; if no symptoms, normal LV function and no arrhythmias allowed to continue with all competitive sports. Follow-up according to individual case.</td>
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recommended before exercise resumption.
Unclear whether myocarditis-related LGE should resolve first.

<table>
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<th>Differences:</th>
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<td>While ACC/AHA scientific statements mention a possible minimum of 3 months of abstinence from sport, European consensus and position papers tend to focus more on 6 months of abstinence. European recommendations included the necessity of absence of symptoms, LV ventricular dysfunction and arrhythmias upon clinical assessment, while ACC/AHA recommendations also consider biomarkers and the possibility of CMR.</td>
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Figure legends
Athlete with current or recent infection

Presence of cardiovascular symptoms
(e.g., Chest Pain, Palpitations, Dizziness, Dyspnea, Syncope, Unexplained performance drop)

- Diagnostic Criteria
  - ECG/Holter/Stress ECG
  - 3T changes in multiple leads
  - PQ depressions, Anhythmia
  - Bio/inflammatory markers
    - Enzyme heart enzymes (Troponin, Creatine kinase), elevated BNP or NT pro-BNP, cytokine inflammation biomarkers
  - Echocardiography
    - Depressed LVFP, RVFP
    - Regional wall motion abnormalities
    - Pericardial effusion

Min. 1 criteria positive

- Endomyocardial biopsy

  Consider treating with immunosuppressants
  treatment depending on EMB results

- Unexplained depressed LVFP

Myocarditis confirmed by LLC evidence of myocardial edema with non-ischemic myocardial injury (LGE) +/- LV dysfunction +/- pericardial effusion

Sports restriction for 3 - 6 months

Follow-up CMR and Stress-ECG after 3 months

- If stable or declining LGE, no arrhythmia and no inflammation
  - Individual case by case recommendation:
    - LGE presence: size and pattern dependent consideration of slow return to exercise over the next 3 - 6 months
    - LGE + reduced LV function: balance slow return to exercise against risk of lack of exercise

- If clinical no inflammation, asymptomatic, normal Stress-ECG and normal CMR without adverse LV/RV remodeling
  - No Sports restriction but consider follow-up

Athlete with clinically suspected myocarditis

CMR risk stratification
- Evaluation of LVEF, RVEF
- Regional wall motion abnormalities
- LGE size, pattern, distribution, location
- T2-w imaging, T2 mapping, EGE
- T1 mapping and ECV
- Pericardial effusion

Normal CMR
- If CMR performed within 7 days of onset of symptoms and strong clinical indicators for myocarditis

Normal CMR and no arrhythmia in Holter or Stress-ECG.
- Normal heart enzymes/inflammation biomarkers subsided to normal

Normal CMR
- CAD ruled out by invasive or noninvasive imaging or clinically low pre-test probability for CAD

Sports restriction until infection resolved/afebrile

No cardiovascular symptoms

Unexplained depressed LVFP

Pericarditis
- Only pericardial effusion, LGE or edema on T2-w imaging only in pericardium, often with CRP elevation and positional pain

Abnormal CMR

Sports restriction for 3 months or until normalization of inflammation

Repeat CMR scan after 2 weeks and sports restriction in meantime

No Sports restriction but consider follow-up
Central Illustration. Proposed diagnostic and treatment algorithm in the assessment of athletes who present with a clinical syndrome of current/recent infection and suspected myocarditis.

BNP = Brain Natriuretic Peptide; CAD = Coronary Artery Disease; ECG = Electrocardiogram; LLC = Lake Louise Criteria; LVEF = Left Ventricular Ejection Fraction; RVEF = Right Ventricular Ejection Fraction; T2-w = T2-weighted;

**Figure 1. Examples of different myocarditis cases in athletes.**

Panel A (CMR scan 10 days after onset of symptoms) represents a case of a 39 years-old recreational athlete who presented with chest pain, increased troponin, ST elevation and depressed LVEF. LGE showed midwall and subepicardial hyperenhancement in the lateral wall (white arrows) with corresponding epicardial myocardial edema (hyperintensity signal) in the T2 weighted images. Treadmill tests revealed ventricular couplet but no cardiac event was reported after a 1-year follow-up. Panel B (CMR scan 3 days after onset of symptoms) shows a soccer player presenting with syncope during exercise. Laboratory testing’s showed no troponin increase but T waves inversion were seen on ECG. The CMR showed no LV hypertrophy, but LV dilation, normal RV dimension and preserved LVEF. There was no myocardial oedema seen on T2 weighted imaging and no LGE present. Acute myocarditis was therefore unlikely. Multiple serial follow-up exercise testing and ECG Holter were normal. Competitive physical activity was resumed with no further cardiac event during a 9-month follow-up. Panel C represents a 49 years old recreational athlete with acute chest pain. EKG showed ST elevation and Troponin I was elevated with 44792 ng/L. Invasive coronary angiogram could exclude a coronary artery disease. The CMR was performed 3 days after symptoms onset and LGE images showed epicardial enhancement on the lateral wall consistent with acute myocarditis. T2-
weighted imaging showed a hyperintensity signal in the lateral wall and T2 mapping depicted increased T2 time in the same region (60 ms) compared to the inferior wall and the septum (36 ms). These findings corresponded to intramyocardial edema. Native T1 mapping native showed increased T1 time in the lateral wall (1466 ms) compared to septum (1280 ms), anterior (1293 ms) and inferior wall (1329 ms).

LV = left ventricular; EF = ejection fraction; ESV = end-systolic volume; EDV = end-diastolic volume; RV = right ventricular; T2-w = T2-weighted

Figure 2. Case of a 62-year-old competitive triathlete who experienced vertigo and syncope of 10 seconds on maximal exertion during cycling.
Electrocardiogram and 24h Holter were normal. Ergometry was normal except ventricular triplets. Coronary arteries were normal on invasive coronary angiography. Echocardiography was normal except mild dilatation of all cardiac chambers. Severe upper respiratory tract infection was remembered two months ago. Laboratory testing including inflammation parameters and cardiac biomarkers were normal. CMR 1 months after the syncope showed large extent of epicardial late gadolinium enhancement (panel A and B, white arrows) anterior and inferior/inferolateral (midventricular) and small amount in the septum seen on the two-chamber,
three chamber and short-axis view. There was no corresponding myocardial edema visible in the T2-weighted imaging (panel C, short axis view shows signal intensity ratio of myocardium compared to skeletal muscle less than 2.0). The diagnosis of myocarditis was made (however, not “acute” at the time of the CMR scan). The athlete was recommended to abstain from strenuous exercise. After implantation of a Reveal recorder, the athlete continued (against the advice of the physician) with high intensity training and experienced a pre-syncope and a ventricular tachycardia of 333/min for 8 seconds (1 months after the CMR was done) recorded on the reveal device (panel D). The recommended beta-blocker was taken infrequently by the athlete and he denied any further electrophysiology study or intra-cardiac defibrillator implantation evaluation. He continued his high intensity trainings (again against the physician advice) and five months after first syncope the athlete died during a strenuous training of a sudden cardiac death.