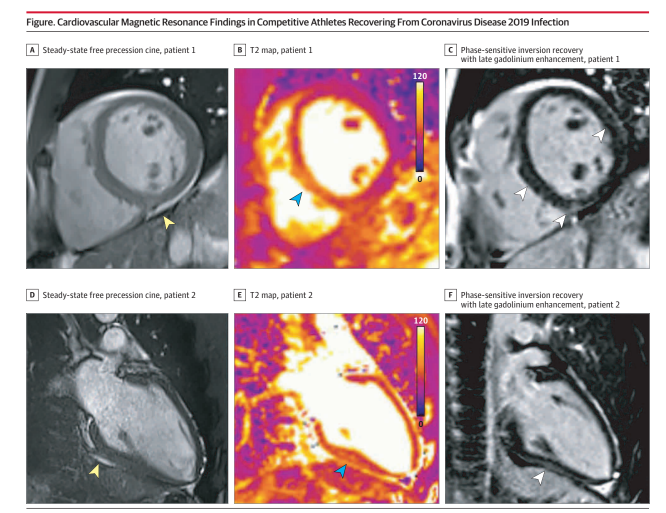
## Letters

## **RESEARCH LETTER**

## Cardiovascular Magnetic Resonance Findings in Competitive Athletes Recovering From COVID-19 Infection

Myocarditis is a significant cause of sudden cardiac death in competitive athletes and can occur with normal ventricular function. Recent studies have raised concerns of myocardial inflammation after recovery from coronavirus disease 2019 (COVID-19), even in asymptomatic or mildly symptomatic patients. Our objective was to investigate the use of cardiac magnetic resonance (CMR) imaging in competitive athletes recovered from COVID-19 to detect myocardial inflammation that would identify high-risk athletes for return to competitive play.

Methods | We performed a comprehensive CMR examination including cine, T1 and T2 mapping, extracellular volume fraction, and late gadolinium enhancement (LGE), on a 1.5-T scanner (Magnetom Sola; Siemens Healthineers) using standardized protocols, <sup>3</sup> in all competitive athletes referred to the sports medicine clinic after testing positive for COVID-19 (reverse transcriptase-polymerase chain reaction) between June and August 2020. The Ohio State University institutional review board approved the study, and informed consent in writing was obtained from participating athletes. Cardiac magnetic resonance imaging was performed after recommended quarantine (11-53 days). Electrocardiogram, serum troponin I, and transthoracic echocardiogram were performed on day of CMR imaging.



A, Cine mid short-axis images showing pericardial effusion indicated by yellow arrowhead. B, T2 map with color overlay mid short-axis showing myocardial edema (elevated T2, 61 milliseconds) indicated by blue arrowhead. C, Short-axis view showing late gadolinium enhancement in the mid inferoseptum, right ventricular insertion point, and mid anterolateral wall indicated by white arrowheads. D, Cine 2-chamber long-axis view showing pericardial effusion indicated by yellow arrowhead. E, T2 map with color overlay myocardial edema (elevated T2, 58 milliseconds) indicated by blue arrowhead. F, Right 2-chamber long-axis view showing epicardial late gadolinium enhancement in the inferior wall indicated by white arrowhead.

Table. Demo	graphic Feat	ures and Echoca	ardiographic and Ca	ardiovascular l	Magnetic Reson	ance Paran	neters in Co	mpetitive Athl	etes Recov	Table. Demographic Features and Echocardiographic and Cardiovascular Magnetic Resonance Parameters in Competitive Athletes Recovering From Coronavirus Disease 2019	irus Disease 2019ª	
			Time CMR performed after	Echocardiog	Echocardiography, mL/m <sup>2</sup>	CMR, %		Native T1		Maximal T7 ms		CMR (undated Lake
Athlete No.	Sex	Symptoms	result, d	LVEDV	RVEDV	LVEF	RVEF	ms ms	ECV, %	(AHA segments)	LGE (pattern/AHA segments)	Louise Criteria)
1	Male	No	21	Not done	Not done	09	49	1034	21	51(9)	Yes (RV insertion; 9)	Normal
2	Male	No	22	51	46	99	59	964	24	48 (9)	Yes (patchy; 6, 8)	Normal
3	Male	No	22	65	09	09	64	953	22	48 (10)	Yes (patchy, 5)	Normal
4	Male	No	15	65	48	59	54	902	20	48 (9)	Yes (linear; 8, 12)	Normal
2	Male	No	17	99	57	55	54	994	24	55 (9)	Yes (epicardial; 3, 9)	Myocarditis
9	Male	Yes	23	73	52	61	62	947	26	63 (3, 9)	Yes (patchy; 3, 9)	Myocarditis
7	Male	Yes	53	99	64	53	52	991	25	49 (7, 9)	Yes (linear, patchy; 8, 9, 12)	Normal
∞	Male	No	20	9/	36	99	53	963	17	51 (10)	No	Normal
6	Male	Yes	18	09	71	99	52	964	24	52 (7)	Yes (patchy; 3, 9)	Normal
10	Male	Yes	11	29	70	61	58	929	25	58 (8, 9)	Yes (patchy; 2, 3, 8, 9)	Myocarditis
11	Male	No	23	57	49	63	09	286	22	53 (7)	No	Normal
12	Male	Yes	28	72	59	20	53	996	28	53 (7, 8)	No	Normal
13	Male	No	28	81	52	33	53	925	25	53 (7, 8)	No	Normal
14	Male	No	11	46	41	65	54	686	24	53 (8)	No	Normal
15	Male	No	48	26	51	59	57	1003	25	53 (7)	Yes (RV insertion; 9)	Normal
16	Female	Yes	23	89	50	64	58	1001	26	52 (8)	No	Normal
17	Female	Yes	23	55	56	57	09	1030	28	48 (10)	No	Normal
18	Female	No	21	53	35	65	99	1008	25	48 (9)	No	Normal
19	Female	Yes	17	09	32	63	57	876	26	53 (8)	No	Normal
20	Female	No	31	62	51	58	59	1002	25	52 (8)	No	Normal
21	Female	Yes	31	52	40	09	09	946	28	53 (8)	No	Normal
22	Female	Yes	30	29	49	59	64	1000	27	52 (8)	Yes (linear; 12)	Normal
23	Female	Yes	30	58	57	57	55	964	26	53 (11)	No	Normal
24	Female	Yes	26	52	49	55	57	1010	30	53 (10)	No	Normal
25	Female	No	31	99	36	99	99	1027	28	50(7)	No	Normal
26	Male	No	12	80	44	09	53	696	21	61(8)	Yes (linear; 8, 9)	Myocarditis
Abbreviation: ECV, extracell	s: AHA, Ameri	can Heart Associa action; EDV, end-	Abbreviations: AHA, American Heart Association; CMR, cardiovascular magnetic resonance imaging: ECV, extracellular volume fraction; EDY, end-diastolic volume; EF, ejection fraction; LGE, late gadolinium	cular magnetic ejection fractio	resonance imagi	ng; Iinium	seg Gar	gmentation usin diovascular ma	ig endocardi. gnetic reson.	al and epicardial contc ance imaging-derived	segmentation using endocardial and epicardial contours at end systole and end diastole per standard protocol Gardiovascular magnetic resonance imaging-derived myocardial T1 and T2 mapping and ECV were done per	per standard protocol.
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segmentation using endocardial and epicardial contours at end systole and end distole per standard protocol. Cardiovascular magnetic resonance imaging-derived myocardial T1 and T2 mapping and ECV were done per standard guidelines. Mean (SD) native T1 less than 999 (3) milliseconds, native T2 of less than 53 milliseconds, and ECV of less than 53 milliseconds, and ECV of less than 29% were considered normal per institutional protocol based on phantom and human volunteer experiments. T2 and LGE were only considered significant if seen in 2 orthogonal planes.

enhancement; LV, left ventricular; RV, right ventricular.

<sup>9</sup> Symptoms refer to symptoms during short-term infection. Echo volumes were calculated by 3-dimensional method. Cardiovascular magnetic resonance imaging-derived left and right ventricular volumes and function were measured from contiguous short-axis cine images using semiautomated software for endocardial

Results | We performed CMR imaging in 26 competitive college athletes (mean [SD] age, 19.5 [1.5] years; 15 male [57.7%]) from the following sports: football, soccer, lacrosse, basketball, and track. No athletes required hospitalization or received COVID-19-specific antiviral therapy. Twelve athletes (26.9%; including 7 female individuals) reported mild symptoms during the short-term infection (sore throat, shortness of breath, myalgias, fever), while others were asymptomatic. There were no diagnostic ST/T wave changes on electrocardiogram, and ventricular volumes and function were within the normal range in all athletes by transthoracic echocardiogram and CMR imaging. No athlete had elevated serum levels of troponin I. Four athletes (15%; all male individuals) had CMR findings consistent with myocarditis based on the presence of 2 main features of the updated Lake Louise Criteria: myocardial edema by elevated T2 signal and myocardial injury by presence of nonischemic LGE (Figure).4 Pericardial effusion was present in 2 athletes with CMR evidence of myocarditis. Two of these 4 athletes with evidence of myocardial inflammation had mild symptoms (shortness of breath), while the other 2 were asymptomatic. Twelve athletes (46%) had LGE (mean of 2 American Heart Association segments), of whom 8 (30.8%) had LGE without concomitant T2 elevation (Table). Mean (SD) T2 in those with suspected myocarditis was 59 (3) milliseconds compared with 51 (2) milliseconds in those without CMR evidence of myocarditis.

Discussion | Of 26 competitive athletes, 4 (15%) had CMR findings suggestive of myocarditis and 8 additional athletes (30.8%) exhibited LGE without T2 elevation suggestive of prior myocardial injury. COVID-19-related myocardial injury in competitive athletes and sports participation remains unclear. Cardiac magnetic resonance imaging has the potential to identify a high-risk cohort for adverse outcomes and may, importantly, risk stratify athletes for safe participation because CMR mapping techniques have a high negative predictive value to rule out myocarditis.4 A recent study by Puntmann et al<sup>2</sup> demonstrated cardiac involvement in a significant number of patients who had recovered from COVID-19. A recent expert consensus article recommended 2-week convalescence followed by no diagnostic cardiac testing if asymptomatic and an electrocardiogram and transthoracic echocardiogram in mildly symptomatic athletes with COVID-19 to return to play for competitive sports. 5 However, emerging knowledge and CMR observations question this recommendation. Cardiac magnetic resonance imaging evidence of myocardial inflammation has been associated with poor outcomes, including myocardial dysfunction and mortality.6 Study limitations include lack of baseline CMR imaging and variable timing of CMR imaging from a positive COVID-19 test result. Athletic cardiac adaptation could be responsible for these abnormalities; however, in this cohort, mean (SD) T2 in those with suspected myocarditis was 59 (3) milliseconds vs 51 (2) milliseconds in those without, favoring pathology. Additionally, the rate of LGE (42%) is higher than in previously described normative populations. To conclude, while long-term follow-up and large studies

including control populations are required to understand CMR changes in competitive athletes, CMR may provide an excellent risk-stratification assessment for myocarditis in athletes who have recovered from COVID-19 to guide safe competitive sports participation.

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Concept and design: Rajpal, Tong, Borchers, Obarski, Simonetti, Daniels. Acquisition, analysis, or interpretation of data: All authors.

Drafting of the manuscript: Rajpal, Tong, Simonetti.

Critical revision of the manuscript for important intellectual content: All authors. Statistical analysis: Rajpal, Tong, Obarski.

Administrative, technical, or material support: Rajpal, Tong, Zareba, Simonetti, Daniels.

Supervision: Rajpal, Tong, Borchers, Daniels.

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