Cardiac injury before and after COVID-19. A longitudinal MRI study

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Background: Recent MRI-based studies have raised great concern about frequent cardiac involvement even in mild or asymptomatic COVID-19. However, while signs of myocardial injury were found in large proportions of patients after COVID-19, all studies published to date lack baseline imaging and are therefore unable to discriminate between pre-existing and COVID-19-induced injury.

Purpose: In this longitudinal study, we aimed to assess the true cardiac impact of COVID-19 based on pre- and post-COVID-19 late gadolinium enhancement (LGE)-MRI.

Methods: A prospective registry of patients with serial LGE-MRIs was screened for patients with documented SARS-COV-2 infection after cardiac LGE-MRI. Eligible patients then received a post-COVID-19 LGE-MRI using the same scanner and sequence as in the pre-COVID-19 MRI. Inversion recovery prepared T1-weighted gradient echo sequences were acquired in sinus rhythm using ECG gating and a free-breathing 3D navigator, 15–20 minutes after administering an intravenous bolus of 0.2 mmol/kg of gadobutrol. A TI scout sequence was used in order to determine the optimal TI that nullified the left ventricular myocardial signal. The presence of LGE was independently assessed qualitatively by two experienced investigators blinded to patient information. For quantitative analyses a 3D-

reconstruction of the left ventricle was performed using ADAS-3D software. LGE was then automatically quantified based on a prespecified signal intensity threshold of \geq 3 SD above the mean of a remote non-enhanced myocardial region.

Results: Pre- and post-COVID LGE-MRI from 31 patients with cardiovascular risk factors that had recovered from mild to moderate COVID-19 (23% hospitalised) were analysed. At a median of 5 months post-COVID-19, LGE-lesions indicative of myocardial injury were encountered in 15 out of 31 patients (48%), which is in line with previous reports. However, intraindividual comparison with the pre-COVID-19 MRI reveiled all of these lesions as pre-existing and thus not COVID-19-related. Quantitative analysis detected no increase in the size of individual LGE-lesions, nor in the global left ventricular LGE-extent. There was no difference in any functional or structural parameter between pre- and post-COVID-19 MRI.

Conclusion: This longitudinal study in a cohort of patients considered at high risk of cardiac involvement, did not find any evidence for COVID-19-induced myocardial injury. The complete absence of de novo LGE lesions in this cohort is reassuring and indicates that cardiac sequelae of COVID-19 are rare and certainly not as common as previously suggested.

LGE post-COVID-19

(transm.) ext

Changes

AHA-segme

Qualitative LGE analysis pre- and post-COVID-19

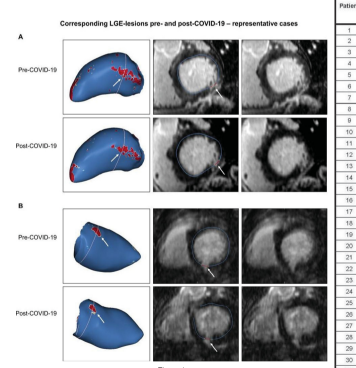
LGE

LGE pre-COVID-19

AHA-segme

(transm.) exte

LGE



/ pattern presen / pattern resentR RVIP RVIP yes yes unchanged n.a. n.a 2: RVI mid: n.a 2: RVI yes yes mid: n.a unchanged 3 yes 2 mid yes 2 mid unchanged 4 no no unchanged 4:2 4; 2 5 yes trans; mid yes trans; mid unchanged 6 yes 8.9; RVIF trans; n.a. 8.9; RVIF trans; n.a. unchanged yes 2, RVIF 2, RVIF yes mid. n.a. yes mid. n.a unchanged no no unchanged 8 unchanged 9 no no 10 yes RVIF RVIP unchanged n.a yes n.a RVIP RVIP unchanged yes n.a yes n.a 12 ves 4 mid yes 4 mid unchanged 13 no no unchanged 14 no по unchanged 15 2,3 mid 2,3 mid yes yes unchanged 5,11; 4,3; RVIP 16 5,11; 4.3; RVIP yes trans; mid; n.a. yes trans; mid; n.a. unchanged unchanged 17 no no 18 no no unchanged unchanged 19 mid mid yes yes 20 7 mid 7 yes yes mid unchanged 21 no no unchanged unchanged 22 no no 23 yes subend yes subendo unchanged 24 RVIP RVIP unchanged yes n.a yes n.a 25 yes RVIP n.a. yes RVIP n.a. unchanged 26 2. RVIE mid' n a 2 RVIE mid n a yes yes unchanged 27 yes 2,3 mid Yes 2,3 mid unchanged 28 unchanged no no 29 RVIP RVIP ves n.a ves n.a unchanged unchanged 30 yes 2 subendo yes 2 subendo 31 ves 3,4 subend 3,4 subendo unchanged

Figure 1

Independent lesions are separated by semicolon, confluent lesions by comma. RVIP = right ventricular insertior point (AHA segments and transmural extent do not apply in RIVP); mid = mid-myocardial; trans = transmural; sub-endocardial.

Table 1