High-resolution Free-breathing late gadolinium enhancement
Cardiovascular magnetic resonance to diagnose myocardial injuries following COVID-19 infection

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Abstract

Purpose: High-resolution free-breathing late gadolinium enhancement (HR-LGE) was shown valuable for the diagnosis of acute coronary syndromes with non-obstructed coronary arteries. The method may be useful to detect COVID-related myocardial injuries but is hampered by prolonged acquisition times. We aimed to introduce an accelerated HR-LGE technique for the diagnosis of COVID-related myocardial injuries.

Method: An undersampled navigator-gated HR-LGE (acquired resolution of 1.25 mm3) sequence combined with advanced patch-based low-rank reconstruction was developed and validated in a phantom and in 23 patients with structural heart disease (test cohort; 15 men; 55 ± 16 years). Twenty patients with laboratory-confirmed COVID-19 infection associated with troponin rise (COVID cohort; 15 men; 46 ± 24 years) prospectively underwent cardiovascular magnetic resonance (CMR) with the proposed sequence in our center. Image sharpness, quality, signal intensity differences and diagnostic value of free-breathing HR-LGE were compared against conventional breath-held low-resolution LGE (LR-LGE, voxel size 1.8x1.4x6mm).

Results: Structures sharpness in the phantom showed no difference with the fully sampled image up to an undersampling factor of x3.8 (P > 0.5). In patients (N = 43), this acceleration allowed for acquisition times of 7min21s ± 1min12s at 1.25 mm3 resolution. Compared with LR-LGE, HR-LGE showed higher image quality (P = 0.03) and comparable signal intensity differences (P > 0.5). In patients with structural heart disease, all LGE-positive segments on LR-LGE were also detected on HR-LGE (80/391) with 21 additional enhanced segments visible only on HR-LGE (101/391, P < 0.001). In 4 patients with COVID-19 history, HR-LGE was definitely positive while LR-LGE was either definitely negative (1 microinfarction and 1 myocarditis) or inconclusive (2 myocarditis).

Abbreviations: HR-LGE, High-resolution late gadolinium enhancement; CMR, Cardiovascular magnetic resonance; LR-LGE, Low-resolution late gadolinium enhancement; LGE, Late gadolinium enhancement; MINOCA, Myocardial infarction with non-obstructed coronary arteries; 3D, Three-dimensional; PCR, Polymerase chain reaction; ECG, Electrocardiogram; LV, Left ventricle; TI, Inversion time; RV, Right ventricular; EF, Ejection fraction; TTE, Transthoracic echocardiography; ROI, Regions-of-interest; BMI, Body mass index.

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Conclusions: Undersampled free-breathing isotropic HR-LGE can detect additional areas of late enhancement as compared to conventional breath-held LR-LGE. In patients with history of COVID-19 infection associated with troponin rise, the method allows for detailed characterization of myocardial injuries in acceptable scan times and without the need for repeated breath holds.

1. Introduction

One major concern related to the recent COVID-19 pandemic lies in the impact of the infection on the myocardium [1]. Several large series have reported high rates of myocardial injuries in hospitalized patients (20–36%), with cardiac involvement being a major factor of adverse outcome [2–4]. Multiple mechanisms can lead to cardiac damage, including type 1 myocardial infarction, type 2 myocardial infarction, stress cardiomyopathy, microvascular ischemia, and myocarditis [5]. Cardiovascular magnetic resonance (CMR) plays a pivotal role in identifying the underlying mechanism in myocardial infarction with non-obstructed coronary arteries (MINOCA) [6], and was thus proposed to detect and characterize COVID-related myocardial injuries. A recent study reported than 49% of patients with history of COVID-19 and troponin rise had positive late gadolinium enhancement (LGE) on CMR [7,8]. The burden that these scars may weigh on the infected population over the long term remains uncertain. Considering the prognostic role of LGE in structural heart diseases and acute coronary syndromes, LGE CMR may play an important role to stratify the risk of future adverse outcomes after COVID-19 infection.

All prior CMR studies in COVID patients have employed conventional LGE techniques acquired during breath holds. While providing good contrast between the injured and healthy myocardial tissue, these techniques have inherently low spatial resolution, which can affect their ability to resolve small myocardial defects (14). In addition, conventional LGE techniques require repeated breath-holds, which can be difficult to perform in COVID-19 patients. High resolution three-dimensional (3D) LGE methods acquired during free-breathing were shown valuable to improve the sensitivity of CMR in the context of MINOCA and to enhance the diagnostic value in atrial fibrosis assessment as well as for the detection of small myocardial scars in ischemic and non-ischemic cardiomyopathies [9–11]. These methods may be useful to detect subtle COVID-related myocardial injuries, particularly in patients with persistent dyspnea, but remain restrained by quite prolonged acquisition times (~8–12 min for anisotropic resolution [9] and ~16 min at 1.4 mm³ isotropic resolution [12,13]). The increasing need for CMR in the context of COVID imposes an optimization of clinical workflows, and accelerated techniques are thus critically needed.

The aim of the present study was to introduce an accelerated free-breathing high-resolution 3D LGE technique to optimize the detection and characterization of COVID-related myocardial injuries.

2. Material and methods

2.1. Study design and patients

The study design consisted of 3 steps. Firstly, the accelerated high resolution free breathing 3D LGE sequence was tested and optimized in a resolution phantom. Secondly, it was validated in a series of patients with known structural heart disease (test group; N = 20), to assess the spectrum of COVID-related myocardial injuries at high spatial resolution. These were consecutive patients prospectively included from May 2020 to November 2020. Inclusion criteria were laboratory confirmed COVID-19 infection associated with troponin rise. Patients requiring mechanical ventilation and those with persistent positive polymerase chain reaction (PCR) testing were not considered for inclusion. The study was approved by an ethics committee, and all patients provided informed consent.

2.2. Whole-heart high-resolution diaphragmatic navigated undersampled 3D LGE

HR-LGE was performed using a free-breathing, three-dimensional, inversion-recovery-prepared, electrocardiogram (ECG)-gated, respiration-navigated gradient-echo pulse sequence (Fig. 1). A SPIR (spectral presaturation with inversion recovery) fat saturation pulse was used to minimize fat-related artefacts. The sequence was accelerated using a 3D variable density Cartesian trajectory [14,15] that samples the $k_x$-$k_y$ phase-encoding plane following approximate spiral interleaves on the Cartesian grid with variable density along each spiral arm (i.e., the k-space center being fully sampled with exponentially increased undersampling towards the periphery). This pseudo-random sampling scheme results in noise-like artefacts in the image domain, which are desired from a reconstruction and compressed-sensing point of view [16]. A spectrally selective crossed-pair navigator preceded each spiral-like interleave to dynamically track the respiratory motion of the heart. An acceptance window of $\pm$ 3 mm was placed in end-expiration.

Imaging parameters are provided in Table 1. As opposed to our previous HR-LGE sequence [9], where 3D data was acquired with mild undersampling factors (GRAPPA x2) and anisotropic resolution (1.25 x 1.25 x 2.5 mm), our proposed isotropic (1.25 mm³) HR-LGE sequence exploits higher undersampling factors (3.8-fold) to achieve clinically feasible (sub-10 min) acquisition times. With such undersampling factor, the image reconstruction pipeline needs to be adapted to compensate for the increased noise due to the ill-posed nature of the reconstruction problem. A patch-based low-rank reconstruction was used to recover high quality LGE images [14]. This reconstruction framework assumes that the 3D HR-LGE data contains a rich amount of redundancy on a local (i.e., within a small patch) and non-local (i.e., between similar patches within a spatial neighbourhood) scales. This redundancy information can be exploited through patch-based matrix representation and singular value truncation to regularize the overall reconstruction problem. The different steps of the reconstruction algorithm are detailed in [14,17]. To assess the full clinical performance of the proposed HR-LGE sequence, the patch-based reconstruction was written in C++ and integrated into the scanner reconstruction software (Image Calculation Environment, Siemens Healthcare, Erlangen, Germany).

2.3. Phantom study

A multi-purpose resolution phantom (Siemens Healthcare, Erlangen, Germany) was imaged with the proposed HR-LGE framework to evaluate the impact of undersampling on image sharpness and resolution. The phantom contains 55 holes with varying diameters in a range of [1.1; 11.5 mm] (Fig. 2). Data was acquired assuming 100% respiratory scan efficiency at a simulated heart rate of 70 beat-per-minute. Different undersampling factors were tested ([1, 2.5, 3.8, 5.2]). The relevant imaging parameters are reported in Table 1. The nominal field-of-view
was chosen to be 23 cm³ for a spatial resolution of 1.25 mm³. Sharpness was measured for each undersampled image at holes' edges as \( \frac{1}{d} \) (in mm⁻¹) with \( d \) being the distance transitioning from 80% to 20% of the intensity profile (given by the minimum and maximum intensities of the profile). Sharpness values were averaged over different positions on the phantom.

### 2.4. Clinical CMR studies

CMR was performed on a 1.5 T clinical scanner (MAGNETOM Aera, Siemens Healthcare, Erlangen, Germany) with a dedicated 32-channel spine coil and an 18-channel body coil. The CMR protocol (Supplementary Fig. 1) comprised Cine SSFP imaging in 2-chamber, 3-chamber, 4-chamber views, and in a stack of contiguous short axis slices encompassing the ventricles. T2, T1 and ECV mapping were also performed in a stack of contiguous short axis slices covering the whole left ventricle (LV).

LR-LGE (reconstructed spatial resolution of \( 1.5 \times 1.5 \times 4.0 \) mm) imaging was performed 10 min after the administration of 0.2 mmol/kg gadoteric acid using a breath-held and inversion recovery-prepared turbo FLASH sequence in 3 stacks of contiguous slices encompassing the ventricles. T2, T1 and ECV mapping were also performed in a stack of contiguous short axis slices covering the whole left ventricle (LV).

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### 2.5. CMR analysis

LV and right ventricular (RV) volumes, ejection fraction (EF) and wall motion abnormalities were analysed by experienced observers (S.S. and H.C.) using a dedicated software (CVI42, Circle Cardiovascular Imaging, Calgary, Canada). The distribution of LGE was categorized as subendocardial, subepicardial, and/or midwall. LGE was considered transmural if involving the entire myocardial thickness on at least one location. The criteria to diagnose myocardial infarction were the presence of subendocardial or transmural LGE [18]. The criteria to diagnose myocarditis were the presence of definite midwall and/or subepicardial LGE in the absence of subendocardial LGE [19]. Arrhythmogenic right ventricular cardiomyopathy was diagnosed according to the Task Force Criteria [20]. The criteria to diagnose takotsubo cardiomyopathy were either a wall motion abnormality involving the entire apical or basal levels in the absence of myocardial LGE, or a similar wall motion abnormality documented on acute transthoracic echocardiography (TTE) in a patient with normal wall motion and negative LGE on CMR [18].

CMR results were interpreted in combination with the clinical history, and after reviewing the results from acute ECG, TTE and biological facts, c) noise artefacts, or d) folding artefacts. For each dataset, the quality was assessed using a three-point ordinal system (1: poor quality with large artefacts, 2: fair quality with moderate artefacts, 3: good quality with small artefacts, 4: excellent quality with no artefacts). For each dataset, the quality was assessed using a four-point ordinal system (1: poor quality with large artefacts, 2: fair quality with moderate artefacts, 3: good quality with small artefacts, 4: excellent quality with no artefacts). For each dataset, the quality was assessed using a three-point ordinal scale (0: presence of subendocardial LGE, 1: presence of subendocardial and subepicardial LGE, 2: presence of subendocardial and transmural LGE, 3: presence of subendocardial, subepicardial, and/or midwall LGE).

### 2.6. LR-LGE versus HR-LGE analysis

LGE images were analysed by 2 readers in consensus (S.S. and A.B., two and seven years' experience in CMR, respectively), with LR-LGE and HR-LGE being reviewed separately 2 months apart, in order to ensure independent analysis of the 2 datasets. Images were reviewed on a clinical PACS-system (Carestream Health, Rochester, New York, USA), where image magnification and windowing could be optimized. Conventional LR-LGE images were reviewed in the four acquired orientations. Owing to its isotropic resolution, the HR-LGE volume could be reviewed in multifocal reformations of any desired orientation, depending on the LGE pattern and distribution.

The overall quality of all LGE images acquired in the test and COVID-19 cohorts was graded using a four-point ordinal system (1: poor quality with large artefacts, 2: fair quality with moderate artefacts, 3: good quality with small artefacts, 4: excellent quality with no artefacts). For scores lower than 4, impaired image quality was classified as a) incomplete myocardial nulling due to sub-optimal TI, b) motion artefacts, c) noise artefacts, or d) folding artefacts. For each dataset, the presence of LGE was assessed using a three-point ordinal scale (0: definitely absent, 1: definitely present, 2: inconclusive) in each of the 17-segment of the American Heart Association model [21]. Therefore, a total of 1462 myocardial segments were analysed in patients.

Regions-of-interest (ROI) placed in the hyperenhancement area, the

| Table 1 | Acquisition parameters for conventional breath-hold low-resolution LGE (LR-LGE) and proposed free-breathing high-resolution LGE (HR-LGE). |
|---------|-------------------------------------------------|
| LR-LGE  | HR-LGE  |
| Repetition time, ms | 3.9 | 5.2 |
| Echo time, ms | 1.7 | 2.5 |
| Flip angle, degrees | 10 | 19 |
| Field of view, mm | \( 360 \times 290 \times 101 \) | \( 360 \times 360 \times 120 \) |
| Acquired resolution, mm | \( 2.0 \times 1.5 \times 7.4 \) | \( 2.5 \times 2.5 \times 1.25 \) |
| Reconstructed resolution, mm | \( 1.5 \times 1.5 \times 4.0 \) | \( 0.63 \times 0.63 \times 0.63 \) |
| Number of slices, range | 11–16 | 96–160 |
| Phase oversampling, % | 15 | 20 |
| Slice oversampling, % | 0 | 16.7 |
| Asymmetric echo | Yes | Yes |
| Acquisition window, ms | 236 | 156 |
| Average number of k-space lines per heartbeat | 30 | 30 |
| Inversion time, ms | 210-300 | 270-350 |
| Trajectory & Acceleration | Cartesian | Cartesian variable density x3.8 |
| GRAPPA x2 | | |
| Trigger pulse, RR interval | 1 | 1 |
| Bandwidth, Hz/pixel | 360 | 255 |
| Breath-hold | Yes | No |

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**Fig. 1.** Schematic overview of investigated free-breathing motion-corrected three-dimensional (3D) whole-heart high-resolution myocardial LGE framework (HR-LGE). A non-selective inversion recovery (IR) pulse is applied immediately after electrocardiogram (ECG) R wave. Image acquisition is gated to account for the diaphragmatic respiratory displacement of the heart. A variable-density Cartesian under-sampling is used to achieve clinically feasible scanning times (undersampling factor of x3.8). The HR-LGE volume is reconstructed with an undersampled patch-based low-rank reconstruction exploiting local patch similarities in the 3D volume. Abbreviations: FAT SAT = fat saturation, \( k_r = \) readout, \( k_p = \) phase encoding, \( k_c = \) slice encoding, TI = inversion time.
remote myocardium, the blood pool and in a background region were extracted for each patient. Injured areas were defined as regions with LGE (based on the two standard deviations segmentation method [22]), while remote areas were defined as regions with no LGE. Mean signal intensity differences and signal ratios were determined by subtracting (respectively dividing) the signal-to-noise ratios (i.e., mean signal level divided by the noise level) of two corresponding tissues.

2.7. Statistical analysis

Results are presented using conventional descriptive statistics. The Shapiro-Wilk test was used to assess for normality. Continuous data are expressed as mean ± standard deviation when following a normal distribution, and as median [interquartile range Q1-Q3] otherwise. Categorical data are expressed as fraction (percentage). Image quality scores of the LR-LGE and HR-LGE datasets were compared using Mann-Whitney U tests. Differences in the detection of LGE segments between the two sequences were tested using the McNemar test. The influence of body mass index (BMI) and heart rate on image quality, presence of artefacts and presence of LGE was assessed using regression analysis. Continuous variables were compared using independent-sample parametric (unpaired Student’s t-test). Statistical tests were 2-tailed. A P value < 0.05 was considered to indicate statistical significance. Statistical analysis was performed using IBM SPSS Statistics (version 26.0).

3. Results

3.1. Phantom study

The total scan times, assuming 100% respiratory gating efficiency, were 15 min 10 sec (x1), 6 min 4 sec (x2.5), 4 min 3 sec (x3.8) and 3 min 3 sec (x5.2) with 1.25 mm³ isotropic resolution. Reconstructed images using the proposed patch-based reconstruction are shown in Fig. 2 in comparison with the reference fully sampled image. Aliasing artifacts can be observed for high undersampling factors (x5) whereas high image quality can be appreciated for undersampling factors up to x3.8, with clear depiction of the phantom structures. Structure sharpness was preserved with the proposed approach (Supplementary Fig. 2) showing no statistical differences with the reference fully sampled image (P > 0.5), as opposed to the standard zero-filling reconstruction (P < 0.01).

3.2. Image characteristics in patients

The mean acquisition time for HR-LGE was 7 min 21 sec ± 1 min 12 sec [min: 3 min 58 sec, max: 9 min 0 sec], with a mean scan efficiency of 42 ± 10 %. The typical image reconstruction time was 1 min.

The overall image quality was significantly higher on HR-LGE compared with LR-LGE (3.65 ± 0.59 vs. 3.09 ± 0.85, P = 0.03). It was graded as excellent (score 4) in 26/41 (63 %) in HR-LGE and 17/41 (41%) in LR-LGE. Reasons for lower image quality were residual motion artefacts (17 vs. 4 patients), sub-optimal TI (7 vs. 6 patients), folding (7 vs. 1 patients), missing slices (3 vs. 0 patients) and residual noise (0 vs. 5 patients) for LR-LGE and HR-LGE, respectively. Nondiagnostic image quality (score 1) was not found in either dataset. On both LR- and HR-LGE, heart rate and BMI did not significantly relate to the image quality, the presence of artefacts, or the presence of LGE (Supplementary Fig. 3).

In terms of contrast, mean signal intensity differences between scar and blood pool and between scar and remote myocardium did not significantly differ between LR and HR-LGE images (P = 0.51 and P = 0.88, respectively). Corresponding signal ratios for LR-LGE relative to HR-LGE were 1.13 for scar/blood pool, 1.01 for scar/remote myocardium, and 1.61 for remote myocardium/blood.

3.3. CMR findings in the test cohort

Baseline characteristics of the test cohort (N = 23; 15 males; mean age 55 ± 16 years) are shown in Table 2. The final diagnosis was ischemic heart disease in 8 (35%) and non-ischemic heart disease in 15 (65%), including 8 (35%) myocarditis, 3 (13%) hypertrophic cardiomyopathy, 3 (13%) dilated cardiomyopathy, and 1 (4%) arrhythmogenic cardiomyopathy with LV involvement. A total of 391 segments were analysed in both datasets (LR-LGE and HR-LGE). Myocardial LGE was found in 23/23 patients (100%) among which 4 (17%) showed transmural patterns, 5 (22%) sub-endocardial, 13 (57%) mid-wall and 1 (4%) sub-epicardial. All enhanced segments that were visible on LR-LGE were also detected on HR-LGE (80/391, 20%) with 21 additional enhanced segments visible only on HR-LGE (101/391, 26%, P < 0.001).
In 3 patients, LGE was categorized as definitely absent on LR-LGE while HR-LGE showed definite mid-wall LGE. In these 3 patients, myocardial LGE was localized in the basal anteroseptal segment, the medial inferoseptal segment and the basal inferolateral and medial inferolateral segments. In 3 other patients, LGE was categorized as inconclusive on LR-LGE, while HR-LGE showed definite LGE (1 mid-wall and 2 transmural patterns). In these 3 patients, myocardial LGE was localized in the basal inferolateral and anteroseptal segments, apical lateral segment, and basal inferolateral, medial inferolateral and apical lateral segments. The benefit of 3D whole-heart isotropic resolution can be appreciated in Fig. 3. The spectrum of myocardial injuries found in the test cohort is illustrated in Fig. 4.

### Table 2
Baseline characteristics of the test cohort (N = 23).

| Patient characteristics | Male gender, N (%) | Age, years | Mean heart rate, beats/min | Mean body mass index, kg/m² |
|-------------------------|--------------------|------------|----------------------------|-----------------------------|
|                         | 15 (65)            | 55 ± 16    | 62 ± 9                     | 25 ± 4                      |
| CMR diagnosis           |                    |            |                            |                             |
| Ischemic cardiomyopathy | 8 (35)             |            |                            |                             |
| Non-ischemic cardiomyopathies | 15 (65)          |            |                            |                             |
| Dilated cardiomyopathy  | 3 (13)             |            |                            |                             |
| Myocarditis, N (%)      | 8 (35)             |            |                            |                             |
| Arrhythmicogenic right ventricular cardiomyopathy | 3 (13) | | | |
| Negative CMR, N (%)     | 1 (4)              |            |                            |                             |

**Cardiac function**

- LVEF, %, mean ± SD: 48.2 ± 16.9
- Reduced LVEF, N (%): 13 (57)
- LV EDV / BSA, mL/m², mean ± SD: 109.4 ± 34.6
- LV ESV / BSA, mL/m², mean ± SD: 58.2 ± 31.5

Note: Data are expressed as mean ± standard deviation unless otherwise specified. Abbreviations: LVEF, left ventricular ejection fraction; EDV, end-diastolic volume; BSA, body surface area; ESV, end-systolic volume; LGE, late gadolinium enhancement.

In 3 patients, LGE was categorized as definitely absent on LR-LGE while HR-LGE showed definite mid-wall LGE. In these 3 patients, myocardial LGE was localized in the basal anteroseptal segment, the medial inferoseptal segment and the basal inferolateral and medial inferolateral segments. In 3 other patients, LGE was categorized as inconclusive on LR-LGE, while HR-LGE showed definite LGE (1 mid-wall and 2 transmural patterns). In these 3 patients, myocardial LGE was localized in the basal inferolateral and anteroseptal segments, apical lateral segment, and basal inferolateral, medial inferolateral and apical lateral segments. The benefit of 3D whole-heart isotropic resolution can be appreciated in Fig. 3. The spectrum of myocardial injuries found in the test cohort is illustrated in Fig. 4.

### 3.4. Characteristics of the COVID-19 cohort

The characteristics of the studied population (N = 20; 15 males; mean age 46 ± 24 years) in the acute phase of COVID-19 are shown in Table 3. Twenty patients were studied, including 2 with history of healed myocardial infarction and 18 with no history of cardiac disease. All patients had been hospitalized, including 11 in an intensive care unit, with 5 requiring mechanical ventilation. COVID-19 diagnosis was based on PCR tests in 16 and serology in 4. The median troponin peak was 305 [Q1:83 - Q3:1134] ng/L. The acute clinical presentation included chest pain in only 7/20 patients. TTE was negative in 11/20. All patients showed either negative or non-specific findings on 12-lead ECG. CMR was performed 54 [32 – 64] days after the onset of symptoms. At the time of the CMR study, 8/20 patients were asymptomatic, while other patients showed a variety of persistent clinical symptoms.

### 3.5. CMR findings in the COVID-19 cohort

Cine imaging showed wall motion abnormalities in 4/20, LV dilatation in 3/20, LVEF impairment in 4/20, and LV thrombus in 1/20 (confirmed on LGE imaging). Edema was found on T2 imaging in 10/20 patients. In 2 patients, HR-LGE imaging could not be completed due to claustrophobia. In these, a definite diagnosis could still be retained based on LR-LGE only (1 takotsubo cardiomyopathy and 1 acute myocarditis).

CMR results are listed in Table 4. Myocardial LGE was found on HR-LGE in 12/18 patients (67%), including 1 (6%) transmural, 3 (17%) sub-endocardial, 4 (22%) mid-wall and 4 (22%) sub-epicardial. All segments with positive LGE on LR-LGE images were also positive on HR-LGE (40/306, 13%) with 8 additional enhanced segments visible only on HR-LGE (48/306, 16%, P < 0.01). In 2 (11%) patients, LGE was definitely absent on LR-LGE while HR-LGE showed definite LGE (one microinfarct and one sub-epicardial LGE). In 2 (11%) other patients, LGE analysis was inconclusive on LR-LGE, whereas HR-LGE showed definite LGE (1 midwall and 1 subepicardial, Supplementary Fig. 4).
The spectrum of myocardial injuries on LGE imaging is illustrated in Figs. 5 and 6. Overall, a definite diagnosis was obtained in 15/20 patients, including myocarditis in 9, multiple microinfarctions in 2, myocardial infarction in 2, takotsubo cardiomyopathy in 2. One additional patient showed signs of acute pericarditis on LGE and T2 imaging but with no signs of myocardial injury. The 4 remaining patients had normal findings on CMR, and no definite diagnosis could be retained. Edema was found on T2 imaging in 8/9 patients with a diagnosis of myocarditis, and in 2/2 patients with a diagnosis of microinfarction. In these 10 patients with edema, the median delay since the onset of symptoms was 28 [16–33] days. The 2 patients diagnosed with takotsubo showed typical wall motion abnormalities on TTE in the acute phase, and negative T2, LGE and cine imaging on the CMR study performed 54 and 20 days after the onset of symptoms. The 4 patients with no definite diagnosis showed a troponin peak of 414 ± 660 ng/mL, normal TTE and ECG findings in the acute phase, and normal CMR results 40 ± 33 days after the onset of symptoms.

4. Discussion

In this study, we demonstrate the feasibility of an accelerated free-breathing high-resolution 3D whole-heart LGE technique for the detection of COVID-related myocardial injuries. The main findings of the present study are that undersampled free-breathing isotropic HR-LGE improves the detection of myocardial defects as compared to conventional breath-held LR-LGE, and that the method allows for detailed characterization of COVID-related myocardial injuries in acceptable scan times.

4.1. Accelerated HR-LGE imaging in the context of COVID

The present study shows that undersampled (x3.8) free-breathing high-resolution isotropic LGE in concert with patch-based reconstruction enables the identification of more injured myocardial segments and improves diagnostic confidence as compared to conventional breath-held LGE methods. This improved sensitivity was equally observed in improvements diagnostic confidence as compared to conventional breath-held LGE methods. The choice of using patch-based low-rank reconstruction over other non-linear reconstruction tools such as compressed sensing [11,31] was motivated by the fact that HR-LGE images contain a rich amount of similar information on a patch scale which can be efficiently exploited though patch-based low-rank reconstruction and consequently lead to better image quality, as previously demonstrated [14,32]. The x3.8 acceleration factor allows for scan times consistently below 10 min for HR-LGE, resulting in a complete CMR protocol duration below 40 min, which is in our opinion compatible with most clinical workflows, and acceptable even in frail COVID patients.

4.2. Spectrum of HR-LGE findings in COVID-19 patients

In the present work, we thought to implement an LGE method specifically suited to the detection of COVID-related myocardial injuries. Because prior reports have shown that cardiac involvement in COVID patients is usually asymptomatic [7] and most often missed on TTE [28], we have anticipated that efficient scan times would be mandatory to allow for the screening of quite large populations on CMR. In addition, prior CMR studies in COVID patients showed that myocardial injuries are most often of small size [8,29,30], justifying efforts to improve the spatial resolution of LGE imaging. Last, the free-breathing approach further alleviated the need for repeated breath holds in COVID patients with impaired respiratory function.

The undersampled free-breathing technique proposed here allows for LGE imaging with near-millimetric isotropic spatial resolution with an overall image quality that favourably compares to the conventional breath held LGE method. The choice of using patch-based low-rank reconstruction over other non-linear reconstruction tools such as compressed sensing [11,31] was motivated by the fact that HR-LGE images contain a rich amount of similar information on a patch scale which can be efficiently exploited through patch-based low-rank reconstruction and consequently lead to better image quality, as previously demonstrated [14,32]. The x3.8 acceleration factor allows for scan times consistently below 10 min for HR-LGE, resulting in a complete CMR protocol duration below 40 min, which is in our opinion compatible with most clinical workflows, and acceptable even in frail COVID patients.
highly prevalent and largely under detected in the infected population, particularly considering that a vast majority of patients is managed outside of hospitals and without laboratory tests to detect potential myocardial injury. Silent myocardial scars are known to be associated with adverse outcomes, exposing to higher risks of ventricular arrhythmia and sudden cardiac death over the long term [34,35]. Thus, there is a critical and urgent need to assess the prevalence of silent scars in the population infected by COVID-19, in order to better appreciate the cardiac risk that the pandemic may weigh on the global population in the near future. In that prospect, the accelerated and high-resolution

### Table 3

| Demographics | Transthoracic echocardiography |
|--------------|-------------------------------|
| Age, years   | 46 ± 24                       |
| Male gender, N (%) | 15 (75)                     |
| LV wall motion abnormalities, N (%) | (15)                  |
| LVEF impairment, N (%) | 3 (15)                     |
| Obesity, N (%) | 7 (35)                        |
| Mean body mass index, kg/m² | 25 ± 7            |
| Smoking, N (%) | 5 (25)                       |
| Hypertension, N (%) | 7 (35)                     |
| Diabetes, N (%) | 2 (10)                       |
| Chronic heart disease, N (%) | 2 (10)                     |
| Acute clinical presentation |                          |
| Fever > 38 °C, N (%) | 13 (65)                     |
| Arhythmia, N (%) | 9 (45)                        |
| Asthenia, N (%) | 14 (70)                      |
| Anosmia or ageusia, N (%) | 7 (35)                     |
| Chest pain, N (%) | 7 (35)                       |
| Dyspnea, N (%) | 10 (50)                      |
| Coughing, N (%) | 12 (60)                      |
| Abdominal symptoms, N (%) | 7 (35)                     |
| Cephalalgia, N (%) | 5 (25)                       |
| COVID-19 diagnosis |                     |
| Chest CT suggestive of COVID-19, N (%) | 11/13 (85)      |
| Positive COVID-19 PCR test, N (%) | 16 (80)             |
| Positive COVID-19 serology, N (%) | 4 (20)                   |
| Laboratory findings |                              |
| C-reactive protein, mg/L, median [IQR] | 88 [59–175]          |
| Fibrinogen, g/L, mean ± SD | 7.1 ± 2.3               |
| NT-proBNP, pg/mL, median [IQR] | 422 [41–479]            |
| Troponin I, ng/mL, median [IQR] | 305 [83–1134]          |

#### Table 4

| CMR findings in the COVID-19 cohort (N = 20). | Timing of the CMR study | Tissue characterization* |
|----------------------------------------------|--------------------------|--------------------------|
| Delay from onset to CMR, days, median [IQR] | 30 [15–58]              | LV native T1, ms, mean ± SD 1028 ± 59 |
| Clinical symptoms at the time of CMR |                           | LV ECV, %, mean ± SD 24.6 ± 2.5 |
| Fever > 38 °C, N (%) | 1 (5)                    | LV T2, ms, mean ± SD 49 ± 4 |
| Arhythmia, N (%) | 2 (10)                   | Focal edema on T2 10 (50) imaging, N (%) |
| Asthenia, N (%) | 8 (40)                   | LGE positive, N (%) 13/20 |
| Anosmia or ageusia, N (%) | 1 (5)                   | Subendocardial LGE, N (%) 3/20 |
| Chest pain, N (%) | 4 (20)                   | Subepicardial or intramural LGE, N (%) 1/20 |
| Dyspnea, N (%) | 2 (10)                   | Transmural LGE, N (%) 1 (5) |
| Coughing, N (%) | 3 (15)                   | LGE burden, N segments, median [IQR] |
| Abdominal symptoms, N (%) | 2 (10)                | RV LGE, N (%) 0/20 |
| Cephalalgia, N (%) | 0 (0)                    | Pericardial LGE, N (%) 1/20 |
| Ventricular function | Final diagnosis | Myocarditis, N (%) 9 (45) |
| LVEDV, mL/m², mean ± SD | 78 ± 19             | Multiple micro-infarction, N (%) 2 (10) |
| LVF%, %, mean ± SD | 56 ± 12              | Myocardial infarction, N (%) 2 (10) |
| LV wall motion abnormality, N (%) | 4 (20)                 | Takotsubo cardiomyopathy, N (%) 2 (10) |
| RVEDV, mL/m², mean ± SD | 80 ± 14             | Pericarditis, N (%) 1 (5) |
| RVF%, %, mean ± SD | 49 ± 9               | Negative CMR, N (%) 4 (20) |
| RV wall motion abnormality, N (%) | 0 (0)                  | Pericardial effusion, N (%) 1 (5) |

* In 2/20 patients LGE results are based on LR-LGE only because HR-LGE was not completed. Abbreviations: CMR, cardiac magnetic resonance; ECV, extracellular volume fraction; EDV, end-diastolic volume index; EF, ejection fraction; LGE, late gadolinium enhancement; LV, left ventricle; IQR, interquartile range Q1-Q3; RV, right ventricle; SD, standard deviation.

LGE technique proposed here may be key to optimize CMR sensitivity.

### 4.3. Study limitations

A limitation of the current study is the unpredictable scan time given by the diaphragmatic navigated technique, where only a fraction of the acquired data is accepted for reconstruction (42 ± 10 % in the present series). Future work should aim at integrating alternate methods to compensate for respiratory motion without impairing scan efficiency, such as image-based navigation [36] or self-navigation [37]. Unfortunately, there was no reference standard for the detection of LGE in this study due to the absence of systematic endomyocardial biopsies to obtain diagnostic confirmation. Endomyocardial biopsy is not part of the routine diagnostic workup in our research centre. Instead, breath-held LR-LGE was used for comparison as this is the reference technique used in our center for assessment of LGE.

One limitation of the study is that the order of the LR-LGE and HR-LGE acquisitions was not randomized, as we could not deviate from our standard clinical protocol in these patients. Even though the relationship between blood and normal myocardium remains in a dynamic equilibrium up to 25 min after contrast injection [38], one may expect an improved scar-to-blood contrast on HR-LGE images as this sequence was always performed after LR-LGE and 15–20 min post injection. Moreover, the prolonged acquisition for HR-LGE can be accompanied by a linear decrease in contrast agent concentration in the blood which could hamper image contrast. Mechanisms based on a gradually increasing TI could be easily integrated into our framework to take into account this effect [39].

Finally, qualitative comparisons with fully sampled HR-LGE dataset.
could not be performed in vivo since it would have come with the caveat of unpractically long (near half an hour) acquisition times and accordingly unfair comparisons due to gadolinium washout and accumulation of motion artefacts. However, phantom experiments supported the use of acceleration factors up to x3.8 to spatially resolve fine structures and recover native image sharpness.

5. Conclusions

Undersampled free-breathing isotropic HR-LGE can detect additional areas of late enhancement as compared to conventional breath-held LR-LGE. In patients with history of COVID-19 infection associated with troponin rise, the method allows for detailed characterization of myocardial injuries in acceptable scan times and without the need for repeated breath holds. Although acute myocarditis is the most common diagnosis, ischemic injuries and takotsubo cardiomyopathy can also be observed. Additional CMR studies are needed to assess the prevalence of occult myocardial scars in the vast majority of infected subjects in whom cardiac troponin was not tested.

Ethics approval and consent to participate

This study was approved by the Biomedical Research Ethics Committee. All patients provided informed consent for participation in this study.

Research Data for this Article

The imaging protocol of the proposed 3D whole-heart HR-LGE sequence as well as the reconstruction framework are available from the corresponding author on reasonable request.

CRediT authorship contribution statement

Aurélien Bustin: Conceptualization, Methodology, Software, Validation, Formal analysis, Data curation, Writing – review & editing.
Soumaya Sridi: Conceptualization, Methodology, Software, Validation, Formal analysis, Data curation, Writing – review & editing.
Pierre Gravinay: Conceptualization, Data curation, Writing – original draft.
Benoit Legghe: . Philippe Gosse: Conceptualization, Data curation, Writing – original draft. Alexandre Ouattara: Conceptualization, Data curation, Writing – original draft. Hadrien Rozé: Conceptualization, Data curation, Writing – original draft. Pierre Coste: Conceptualization, Data curation, Writing – original draft. Edouard Gerbaud: Conceptualization, Data curation, Writing – original draft. Arnaud Desclaux: Conceptualization, Data curation, Writing – original draft. Alexandre Boyer: Conceptualization, Data curation, Writing – original draft. Didier Gruson: Conceptualization, Data curation, Writing – original draft. Nahema Issa: Conceptualization, Data curation, Writing – original draft. Michel Montaudon: Conceptualization, Data curation, Writing – original draft. Francois Laurent: Conceptualization, Data curation, Writing – original draft. Fabrice Bonnet: Conceptualization, Data curation, Writing – original draft.

Fig. 5. Image quality comparisons between conventional breath-held low-resolution LGE (LR-LGE) and the proposed free-breathing high-resolution LGE (HR-LGE, reformatted images) sequence in four patients with laboratory-confirmed COVID-19 infection associated with troponin rise. The final diagnoses were negative CMR (top left), myocardial infarction (bottom left), acute myocarditis (top right and bottom right). Yellow arrows indicate sites of LGE.
Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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