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1	Evaluation of a real-time MRI-guided electrophysiology
2	system for structural and electrophysiological
3	ventricular tachycardia substrate assessment
4 5 6	Mukherjee et al. Real-time MRI-guided electrophysiology
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63	Condensed abstract			
64 65	A real-time MRI-guided electrophysiology (MR-EP) system was used to assess structural and			
66	electrophysiological substrate in a porcine ischaemia-reperfusion model. There was a			
67	moderate correlation between regions of low voltage and delayed conduction identified using			
68	the MR-EP system and late gadolinium enhancement (LGE).			
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107 What's new?

108	•	Endocardial voltage mapping and limited assessments of slow conduction were
109		feasible in a porcine ischaemia-reperfusion model using a novel real-time MRI-
110		guided electrophysiology system (MR-EP)
111	•	Using conventional bipolar voltage thresholds, there was moderate sensitivity in the
112		ability of voltage mapping with the MR-EP system to identify regions of late
113		gadolinium enhancement (LGE)
114	•	An improved sensitivity for LGE detection may be achieved using higher normal
115		bipolar voltage cut-offs with the MR-EP system and respective MR-compatible
116		catheter
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133 Background:

134 Potential advantages of real-time magnetic resonance imaging-guided electrophysiology

135 (MR-EP) include contemporaneous 3D substrate assessment at the time of intervention,

136 improved procedural guidance and ablation lesion assessment.

137

138 **Objective:**

139 We evaluated a novel real-time MR-EP system to perform endocardial voltage mapping and

140 assessment of delayed conduction in a porcine ischaemia-reperfusion model.

141

142 Methods

Sites of low voltage and slow conduction identified using the system were registered and
compared to regions of late gadolinium enhancement (LGE) on MRI. The Sorensen-Dice
similarity coefficient (DSC) between LGE scar maps and voltage maps was computed on a
nodal basis.

147

148 **Results**

149 A total of 445 electrograms were recorded in sinus rhythm (range: 30-186) using the MR-EP system including 138 electrograms from LGE regions. Pacing captured at 103 sites; 47 150 151 (45.6%) sites had a stimulus-to-QRS (S-QRS) delay of \geq 40ms. Using conventional (0.5mV-1.5mV) bipolar voltage thresholds, the sensitivity and specificity of voltage mapping using 152 the MR-EP system to identify MR-derived LGE was 57% and 96% respectively. Voltage 153 154 mapping had a better predictive ability in detecting LGE compared to S-QRS measurements 155 using this system (area under curve: 0.907 vs 0.840). Using an electrical threshold of 1.5mV to define abnormal myocardium, the total DSC, scar DSC and normal myocardium DSC 156

157	between voltage maps and LGE scar maps was 79.0% \pm 6.0%, 35.0% \pm 10.1% and 90.4% \pm
158	8.6% respectively.
159	
160	Conclusions:
161	Low voltage zones and regions of delayed conduction determined using a real-time MR-EP
162	system are moderately associated with LGE areas identified on MRI.
163	
164	Keywords:
165	Real-time, magnetic resonance imaging, electroanatomic mapping, substrate, ventricular
166	tachycardia, late gadolinium enhancement
167	
168	Abstract word count - 237
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192 Introduction

There is growing interest in the use of real-time magnetic resonance imaging-guided
electrophysiology (MR-EP) to treat patients with cardiac arrhythmias.^{1,2} Potential advantages
of MR-EP procedures include soft tissue visualisation with a high contrast-to-noise ratio,
improved assessment of arrhythmia structural substrate using late gadolinium enhancement
(LGE) scar imaging, navigation of catheters using dedicated tracking techniques, online
monitoring of ablation lesion formation and an evaluation of anatomic and physiologic
changes during mapping and lesion delivery.³

200

201 Although most preliminary real-time MR-EP studies have been performed in the atria, where significant technical challenges remain for accurate substrate evaluation,^{1,2,4} MRI is the gold 202 203 standard imaging modality for assessment of ventricular function and scar burden.⁵ 204 Combined MR-EP techniques could offer synergistic benefits for the evaluation and ablation 205 of ventricular tachycardia (VT) substrate. Previous studies using conventional systems and 206 image integration where the association between electrical substrate for VT and MRI-derived 207 scar have been investigated invariably report registration errors on a scale between 3.8 -4.3mm^{6,7} which could be a significant source of mis-match.⁸ Real-time MR-EP enables 208 209 image registration to be performed within a single imaging modality, acquire imaging and 210 electrical data in the same coordinate system and minimise translational changes due to beat-211 to-beat cardiac motion and respiratory motion.

212

213 In this study, we describe the ability of a novel real-time MR-EP system to perform

endocardial voltage mapping and limited assessments of delayed conduction in a porcine

215 ischaemia-reperfusion model taking advantage of custom technical developments in a second

216 generation MR-compatible catheter and a dedicated prototype image-guidance platform for

217	interventional procedures. We hypothesised that with the minimisation of registration errors
218	and translational changes expected using a real-time MR-EP platform, an improved
219	association between structural and electrophysiological substrate may be expected.
220	
221	Methods
222	Animal model and infarct preparation:
223	
224	The research protocol was approved by the local institutional review board and complied
225	with French law on animal experiments and the Guiding Principles for the Care and Use of
226	Laboratory Animals published by the National Institutes of Health (8th Edition, National
227	Academies Press, 2011). The research was performed at the Institut de Chirurgie Guidée par

228 l'image (IHU), Strasbourg, France. Seven male domestic pigs (weight - 35.7 ± 5 kg; 2

healthy, 5 post infarction) were treated with 800mg amiodarone, twice daily for 4 days prior

to and following an infarct procedure and/or imaging and electrophysiology studies. A

231 closed-chest model of myocardial infarction was used as previously described.⁹ (See

supplementary data for detailed methods).

233

All animals underwent a MRI scan for substrate assessment 6 weeks after infarct on a 1.5T scanner (MAGNETOM, Aera, Siemens Healthcare, Erlangen, Germany). Each animal was sedated, intubated and mechanically ventilated as per the infarct procedure for all imaging studies. A 3D ECG-triggered whole heart bSSFP MRI dataset was acquired to enable manual segmentations of cardiac chambers (transverse slice orientation, AP phase encoding, 256 x 256 in-plane matrix size, TR/TE/ α = 3.7ms/1.64ms/90°, voxel size = 1.25x1.25x2.5mm³,

²³⁴ *Imaging study*

²³⁵

242	bandwidth = 895Hx/Px, GRAPPA factor = 2). For scar imaging, contrast was administered
243	(Gadovist, Bayer, Germany) at a dose of 0.2mmol/kg. High-resolution 3D late gadolinium
244	enhancement (LGE) imaging was performed using a free-breathing, respiratory navigator and
245	ECG-gated (in diastole) inversion recovery, b-SSFP sequence ((TR/TE/ α =3.45ms/1.5ms/90°,
246	FOV=339×264×100mm ³ , voxel size=1.2×1.2×1.2mm ³ , bandwidth=895Hz/Px, GRAPPA
247	factor=2, 2RR acquisition). The LGE sequence was run 10-15 minutes after administration of
248	contrast. Based on the LGE-MRI, scar was manually segmented using a version of the
249	Medical Imaging Interaction Toolkit (MITK, Heidelberg, Germany) with the full-width-half-
250	maximum (FWHM) threshold used to define scar and help guide electroanatomic mapping
251	(EAM) during the subsequent procedure.
252	
253	iCMR image guidance platform
254	
255	A custom interventional cardiovascular magnetic resonance (iCMR) image guidance
256	platform (Siemens Healthcare, Erlangen, Germany) was used in this study (Figures 1 and 2).
257	The application has the ability to load volumetric data from MRI scans, display multi-plane
258	reconstructions (MPR) in 3 orthogonal planes and transfer segmentations of cardiac
259	chambers derived from previous imaging or imaging acquired at the time of the EAM

260 procedure. An automatic segmentation tool is incorporated within the software to ensure

rapid image processing. During the MR-EP procedure, the MPR slices on the iCMR

application can follow the tip of the actively tracked catheter to display 3D location of thecatheter within the segmentations of the cardiac chambers as well as on the MPR images

264 (Figure 2). The position of the actively tracked catheter is displayed following the

265 implementation of a temporal smoothing algorithm that limits its excursion due to cardiac

266 motion.

267 The software allows the imaging operator to start/stop sequences remote from the scanner 268 console and configure parameters of each sequence on the MRI scanner. A MR-compatible 269 foot-switch is also available as part of the application to start or pause an interactive imaging 270 sequence that the electrophysiologist can operate (e.g. to use MPRs to navigate the catheter to 271 a region of interest). The mapping interface of the application allows for changes to the rendering style or colour of a loaded segmentation, as well as place markers in regions of 272 273 interest (e.g. to highlight EGMs or mark sites of ablation). The iCMR application 274 communicates directly with the Advantage EP Recording system to display recorded 275 activation times and voltage amplitudes. Colour interpolation is used to display this data 276 which is computed by a relaxation algorithm that takes the values on the mapping points as 277 the fixed boundary condition and then performs a linear interpolation on the segmentation 278 surface between these mapping points. These features enable the system to closely mimic that 279 of a clinical EAM system whilst having additional capabilities to utilise imaging data for procedure guidance. 280

281

282 Real-time MRI-guided electrophysiology procedure

283

Vascular access was obtained via the femoral artery and vein under ultrasound guidance (9Fr 284 285 or 10Fr introducer sheath) followed by administration of 100 units/kg of intravenous heparin. 286 All EAM studies were performed inside the MRI scanner without the use of fluoroscopy at 287 any point. The left ventricle (LV), right ventricle (RV), left atrium (LA), right atrium (RA) and aorta were manually segmented from the 3D ECG-triggered whole heart bSSFP MRI 288 289 dataset using the MITK-based platform. Image processing was performed during a 45-minute 290 window following the completion of imaging studies and prior to the start of EAM. During 291 this time, each animal remained inside the scanner in order to minimise translational changes

due to subject movement between imaging and mapping. The 3D shells of each chamber
were imported into the iCMR guidance platform and displayed using the 3D-whole heart
dataset to act as a 'road-map' for mapping studies. The 3D segmentation of scar from the
LGE-MRI was also imported into the iCMR guidance platform and overlaid onto the 3D
shell for the LV chamber.

297

298 A custom 9Fr, MR-compatible steerable catheter with a single gold 3.5mm tip and ring 299 bipolar electrode (3.5mm inter-electrode spacing) and six circumferential open irrigation 300 ports (Vision-MR, Imricor, Burnsville, MN, USA) was advanced into the LV cavity via 301 retrograde aortic access. A number of modifications were implemented to the MR-compatible catheter from previous versions used in the atria^{1,2} to enable manipulation in the left ventricle 302 303 (Supplementary Data). These changes enabled improved torque transfer within the ventricle, 304 manoeuvrability and consistency of shape following deflection. The MR-compatible catheter has 2 solenoid micro-coils located 2mm and 11mm proximal to the ring electrode that 305 306 enabled the location and orientation of the catheter to be detected in 3D space using a 307 dedicated MRI active tracking sequence. A custom-built MR-EP recording system (Advantage-MR, Imricor, Burnsville, MN, USA) consisting of a digital amplifier, stimulator 308 309 and host workstation was used to record, display and analyse intra-cardiac electrograms as previously described.¹⁰ A patient monitoring system suitable for use in the MRI environment 310 311 (Invivo, Gainesville, Florida) was used to monitor a single lead ECG and invasive arterial 312 blood pressure throughout the study.

313

314

315 Active catheter tracking of MR-compatible catheter

317 In order to accurately detect the location and orientation of the mapping catheter in 3D space, a dedicated active tracking sequence was used as described previously.¹⁰ Briefly, the X, Y, Z 318 coordinates of the catheter micro-coils were determined using the custom active tracking 319 320 sequence, which was optionally interleaved with a fast balanced steady state free precession 321 (bSSFP) imaging sequence automatically following the current catheter position. The active tracking sequence comprised three non-selective projection acquisitions along the respective 322 323 axis. A dynamic imaging coil detuning approach and pre-spoiler were applied to avoid potential background noise, i.e. coil coupling and residual signal effects. Based on the 324 325 acquired projections, the corresponding signal peaks were detected with a dynamic template-326 matching algorithm, which used the initial projections to calculate a template per coil and axis. The template was continuously updated with each new projection fulfilling a minimal 327 peak-to-noise ratio to adapt to the changing shape of the projections while manoeuvring the 328 329 catheter. The detected positions were fed back to both the iCMR platform (Siemens 330 Healthcare) and the MRI scanner to update the rendered catheter position/orientation and 331 imaging plane location respectively.¹⁰

332

333 Intra-cardiac Electrogram recording and characterisation

334

Activation and voltage data were acquired during sinus rhythm. For each sampling point, the time delay (LAT) from a fixed intra-cardiac reference point to the initial deflection of the local LV electrogram was measured manually on the EP recording system and data transferred to the iCMR image guidance platform. Similarly, the peak-to-peak voltage amplitude was also measured manually and transmitted to the guidance platform (Figure 1). Both datasets were used to generate colour-coded activation and voltage maps on the iCMR platform. Areas of focused mapping were based on the location of LGE-derived scar. In order 342 to avoid EGM artifacts due to poor catheter-tissue contact, at least 2 consecutive EGMs had 343 to have the same morphology prior to acceptance of each mapping point. Regions of abnormal myocardium were defined as areas with a bipolar voltage threshold <1.5mV.¹¹ 344 345 EGMs were reviewed off-line at a sweep speed of 100mm/s. After acquisition of activation 346 and voltage maps, the LV catheter was used to pace during stable sinus rhythm (10mA, 3ms, cycle length 10% shorter than sinus cycle length) from sites of normal myocardium and scar. 347 348 The time from the stimulus artefact to the surface QRS onset was used to distinguish regions of normal and delayed conduction. Following confirmation of capture, the time duration 349 350 between the stimulus artefact to QRS onset was recorded. The MR-compatible catheter was 351 sequentially manoeuvred to sites within normal myocardium and scar using active catheter tracking to generate a colour-coded map of stimulus-QRS duration times (S-QRS). Sites with 352 353 a S-QRS >40ms during pace-mapping in sinus rhythm were considered regions of slow conduction as previously described.¹² Following completion of the MR-EP procedure, pigs 354 were euthanised with potassium chloride and hearts were rapidly dissected for gross 355 356 pathological examination. Hearts were photographed with areas of ischaemic scar delineated. 357

358 Image registration, scar segmentation and comparison to voltage maps

359

The LGE-MRI imaging was registered to the 3D whole heart MRI datasets using a pointbased (landmark) rigid registration to guide EAM. Points were selected within the RV, LV and LA blood pools of each image dataset. Registration was performed on the Medical Imaging Interaction Toolkit (MITK) [https://doi.org/10.1016/j.media.2005.04.005]. Scar was segmented on the LGE-MRI using the FWHM method to normalise signal intensity relative to maximum myocardial signal intensity. First, the LV wall was manually segmented using a custom version of MITK. This was performed using the 'Paint Tool' on the MITK-based platform to derive the endocardial and epicardial border on a slice-by-slice basis with 3D
interpolation to minimise discontinuities between slices. Then, the maximum signal intensity
within the LV wall was computed and the voxels with signal intensity above 50% of the
maximum intensity (FWHM) were labelled as scar.

371

372 To compare the scar segmentation with regions of low voltage, the scar segmentation was 373 mapped onto the voltage map surface mesh. This was achieved in two steps. First, the scar segmentation image was rotated and translated so that it was aligned with the surface mesh. 374 375 Second, the scar points were mapped onto the surface mesh using the iterative closest point 376 (ICP) method. In addition, the voltage map was converted to a binary map of scar (1) and normal tissue (0). In this ischaemia-reperfusion model, scar has been noted to be transmural 377 in the majority of myocardial segments with LGE.⁹ The Sorensen-Dice similarity coefficient 378 379 (DSC) between the two binary maps was then computed on a nodal basis for all regions, scar regions only and regions of normal myocardium. The DSC between LGE scar maps and 380 381 voltage maps following thresholding at different cut-offs (0.5mV-3.5mV) was also derived.

382

383 Statistical analysis

384

Data analysis was performed using GraphPad Prism version 7.0 (GraphPad Software, CA,
USA) or SPSS v24.0 (IBM Corp. Armonk, NY, USA). Continuous data are represented as
mean ± SD and compared using the Student's two-tailed T-test. A 2-sided p value <0.05 was
considered statistically significant. For assessment of the accuracy of the MR-EP system to
correctly identify scar and delayed conduction, the location of LGE-derived scar was taken as
the 'gold standard' of structural substrate. The sensitivity, specificity, positive predictive
value and negative predictive value of low voltage points and S-QRS times using the MR-EP

392 system to identify LGE-scar was assessed and used to derive receiver operator characteristic393 (ROC) curves.

394

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All pigs that underwent a LAD infarct developed antero-septal scar which was visualised on
the LGE images (mean scar volume - 6.80 ± 0.88ml) - Supplementary data. There was no
LGE present in healthy pigs that did not undergo the LAD infarct procedure.

399

400 *Real-time MRI-guided electroanatomical mapping*

401

402 Segmentations of scar from the LGE-MRI were displayed on the iCMR image-guidance platform as coloured shells to guide EAM (Figure 2). 445 EGMs (range 30-186) were 403 404 recorded from all animals in sinus rhythm (including 138 EGMs from regions located within the LGE scar segmentation). Using the MRI-derived LGE segmentation to differentiate 405 406 between normal myocardium and scar, the mean signal-to-noise ratio (SNR) of EGMs within normal tissue and scar was 44.78 ± 21.91 and 11.67 ± 6.99 respectively (p<0.0001) (Figure 407 408 3). Pacing captured at 103 sites whilst 10 sites which were all in regions of LGE-derived scar 409 did not capture; 56 (54.4%) sites had S-QRS delay \leq 40ms, 47 (45.6%) sites had a delay of 410 \geq 40ms whilst 15 (14.5%) had a delay \geq 80ms. Representative examples of voltage and S-QRS 411 maps obtained using the system are shown in Figure 4. 412 Relationship between MRI-derived scar, voltage and delayed conduction 413 414 Using conventional (0.5mV-1.5mV) bipolar voltage thresholds, the sensitivity and specificity 415

416 of voltage mapping using the MR-EP system to identify MR-derived LGE was 57% and 96%

417	respectively (ROC area under curve = 0.907 ; p< 0.0001). A S-QRS threshold of >40ms using
418	this system resulted in a sensitivity of 76% and specificity of 73% to identify MR-derived
419	LGE (ROC area under curve = 0.840; p<0.0001) - Figure 5. At a threshold of 1.5mV to
420	define abnormal myocardium, the positive predictive value (PPV) and negative predictive
421	value (NPV) of voltage mapping to identify LGE was 86% and 83% respectively. At a
422	threshold of 40ms, the PPV and NPV of S-QRS time using the system to identify LGE was
423	73% and 79% respectively (Figure 5).
424	
425	There was a moderate relationship between low voltage regions in the LV endocardium and
426	LGE-derived scar mapped onto the endocardial surface mesh (Figure 6). At a voltage
427	threshold of 1.5mV, mean DSC across all nodes was 79.0% \pm 6.0%, whilst mean DSC within
428	scar regions only was 35.0% \pm 10.1% and 90.4% \pm 8.6% in normal myocardium regions
429	only. An improvement in DSC within scar regions was observed using a higher voltage cut-
430	off of 2.0mV and 2.5mV (47.3 \pm 9.9% and 60.2 \pm 22.4%) at the expense of reduced
431	agreement across regions of normal myocardium (Figure 7).
432	
433	Discussion
434	
435	This study shows that the prototype real-time MR-EP system can be used to guide catheters
436	to regions of scar using active catheter tracking and to distinguish regions of low voltage and
437	delayed conduction from healthy myocardium. There is a moderate relationship between low
438	voltage and LGE scar using conventional bipolar voltage thresholds. An improved sensitivity
439	for LGE detection may be achieved using higher bipolar voltage cut-offs with this system.

441 The relationship between local EGM amplitude and scar is complex, in part due to the dependence of voltage on infarct size, heterogeneity and transmurality.¹³ Conventional 442 bipolar voltage thresholds for scar detection may lack sensitivity to fully detect scar as 443 444 variations in inter-electrode spacing and recording electrode size may affect the representation of EGMs.¹¹ Furthermore, although LGE-MRI is the current gold standard for 445 visualisation of ventricular scar post myocardial infarction, the limited spatial resolution of *in* 446 447 vivo LGE-MRI can result in partial volume effects and limit the specificity of scar characterisation.¹³ Increasing mapping resolution using multi-electrode catheters may also 448 449 result in detection of a smaller area of low bipolar voltage as each data point represents a smaller tissue area with less far-field contamination¹⁴. The use of multi-electrode catheters 450 451 could improve the correlation between EAM and imaging as has been shown in a randomised study.¹⁵ An additional source of discrepancy when correlating EAM and pre-procedural 452 453 imaging is registration error due to translational changes (patient movement, cardiac or respiratory motion) or changes in volume, orientation or rhythm of the heart between time of 454 imaging and EAM.¹⁶ 455

456

The real-time MR-EP system minimises registration error through registration of electrical 457 458 and structural data within a single imaging modality with the same coordinate system. The 459 3D whole heart sequence used for chamber segmentation was acquired during the same phase 460 of the cardiac cycle as the 3D LGE to minimise translational changes due to beat-to-beat 461 cardiac motion. Furthermore, both sequences were performed when animals were under general anaesthesia with reduced variability in respiratory motion thereby minimising 462 463 translational changes due to respiratory motion. Compared to image integration approaches, 464 where positional errors are introduced when registering catheter position to pre-procedural imaging, the MR-EP system tracks catheter position directly using a dedicated tracking 465

466 sequence that is acquired in the same coordinate system as the 3D whole heart and LGE 467 scans. The main sources of error with the MR-EP system include within scan registration error and catheter tip displacement on the 3D shell with the active tracking sequence. In a 468 469 cohort of conscious patients scanned with an angiography sequence to create an endocardial 470 mask and a 3D LGE acquisition, the within scan translation error was noted to be 1.9 \pm 1.6mm with a rotation error of $0.62 \pm 0.41^{\circ}$.¹⁷ This is, however, likely to overestimate within 471 scar error with the MR-EP system where translational movements were minimal as animals 472 were under general anaesthesia. Using ex-vivo technical validation, the average tip 473 474 displacement of the actively tracked catheter using the MR-EP system was measured as 0.90 ± 0.58 mm along the axis of the catheter¹ and is likely to be the best estimate of error with this 475 476 set-up.

477

478 In this study, we show that despite the minimisation of registration and translational errors, 479 the relationship between scar delineated using a custom MR-compatible catheter and high resolution isotropic LGE imaging (1.2mm³) remains moderate when using standard voltage 480 481 thresholds. An improvement in scar concordance with this system can be achieved using a higher normal bipolar voltage cut-off. Some investigators have found that abnormal 482 potentials targeted for ablation may be present in tissue classified as 'normal' (>1.5mV) 483 484 voltage and manual adjustment of bipolar voltage thresholds to higher cut-off values may identify more confluent scar regions incorporating all abnormal signals.¹⁸ Regions of slow 485 486 conduction could also be present in tissue of normal bipolar voltage and unmasked during extrastimulus pacing.¹⁹ 487

488

Although the majority of real-time MR-EP studies published previously have focused on theatria, the full potential of substrate and lesion assessment afforded by such systems is likely

to be realised in the context of VT ablation. There are limited data available evaluating realtime MR-EP systems in the ventricle.^{20,21,22} Our study builds on previous work to characterise
the relationship between LGE-derived scar and electrophysiological measurements of low
voltage and delayed conduction inside a MRI scanner.

495

496 Currently, limited visualisation of soft tissue structures is possible in the electrophysiology 497 laboratory with the use of intra-cardiac ultrasound (ICE), however MRI offers an improved contrast-to-noise ratio and ability to acquire 3D whole heart images or 2D slices in any 498 499 imaging plane. Furthermore, tissue characterisation techniques such as LGE can be used to 500 identify arrhythmogenic substrate whilst dedicated sequences can be used to monitor tissue 501 temperature during ablation and provide a real-time method of calculating lethal thermal 502 dose.¹⁰ The novel MR-EP system described is capable of visualising the location and 503 orientation of catheters relative to soft tissue, assess scar with MRI at the time of EAM, 504 enable rapid segmentation and registration of cardiac chambers and potentially monitor formation of ablation lesions.¹⁰ These features of the MR-EP system could offer an 505 506 alternative to conventional fluoroscopy-guided or ICE-guided procedures and improve catheter navigation, delivery of therapy and assess anatomical and physiological changes 507 508 during VT ablation with the potential to reduce risks and improve outcomes.

509

A number of technical developments are required prior to the realisation of real-time MRIguided VT ablation. The development of a MRI-compatible defibrillation system will be a
prerequisite prior to any clinical studies and prototypes are currently under evaluation.²³
Current surface ECG monitoring systems inside a MRI scanner are limited to 4-6 surface
electrodes; in order to aid the diagnostic electrophysiology requirements of VT ablation,
robust 12-lead ECG systems are required. Although high-fidelity 12-lead ECG recordings are

516 possible,²⁴ the impact of magneto-hydrodynamic effects and gradient switching-induced 517 voltages within the MRI scanner can still corrupt ECG signals. There is currently a limited 518 availability of MR-compatible devices; the development of MR-compatible multi-electrode 519 catheters with similar capabilities to their conventional counterparts will accelerate progress 520 in the electrophysiological assessment of substrate inside the scanner.²⁵

521

522 Limitations

523 There are several important limitations to this study. We did not define the bipolar voltage 524 threshold that best correlates to histological scar using the MR-compatible catheter - rather 525 two indirect methods of scar assessment were compared to each other. During assessment of S-QRS intervals to assess slow conduction, a single ECG lead was used to derive 526 527 measurements due to the lack of availability of a MRI-compatible 12-lead ECG; as a result, 528 no assessment of QRS morphology using a 12-lead ECG was performed during pacing. These measurements should therefore be interpreted with caution as we could not account for 529 530 local latency although this would be expected to be minimal at the pacing cycle length used. 531 Furthermore, the technique of S-QRS measurements may have limited sensitivity for the detection of regions of myocardium with slow conduction compared to an approach 532 analysing the evoked response to extrastimuli.¹⁹ In this model, haemodynamic compromise 533 534 and death of the animal was inevitable if VT was induced. As there was no means to 535 defibrillate the animal inside the scanner, we deliberately avoided the induction of VT which 536 in turn precluded entrainment mapping. The MR-EP system used in this study consisted of a single electrode catheter and required manual annotation of activation times and voltages for 537 538 each point on the EP recording system resulting in substantially lower mapping densities than with contemporary EAM systems. This could have lowered the precision of the sensitivity 539 and specificity measures reported in the study. The development of automated mapping 540

relationship between electrophysiological substrate and MR-derived substrate.

543

544 Conclusions

There is a moderate association between low voltage regions and sites of altered conduction determined using a novel real-time MR-EP system with scar derived from LGE-MRI. An improved sensitivity for LGE detection could be achieved using a higher normal voltage cutoff with this system and the respective catheter. Further technical developments in MRcompatible devices will accelerate progress towards real-time MRI-guided VT ablation.

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562 **References**

¹ Chubb H, Harrison JL, Weiss S, Krueger S, Koken P, Bloch LO et al. Development, pre-

564 clinical validation and clinical translation of a cardiac magnetic resonance-electrophysiology

565	system with active catheter tracking for ablation of cardiac arrhythmia. Journal of the
566	American College of Cardiology: Clinical Electrophysiology 2017: 3; 89-103.
567	
568	² Hilbert S, Sommer P, Gutberlet M, Gaspar T, Foldyna B, Piorkowski C et al. Real-time
569	magnetic resonance-guided ablation of typical right atrial flutter using a combination of
570	active catheter tracking and passive catheter visualisation in man: initial results from a
571	consecutive patient series. Europace 2016: 18; 572-7.
572	
573	³ Mukherjee RK, Whitaker J, Williams SE, Razavi R, O'Neill MD. Magnetic resonance
574	imaging guidance for the optimization of ventricular tachycardia ablation. Europace 2018:
575	20; 1721-32.
576	
577	⁴ Paetsch I, Sommer P, Jahnke C, Hilbert S, Loebe S, Schoene K et al. Clinical workflow and
578	applicability of electrophysiological cardiovascular magnetic resonance-guided
579	radiofrequency ablation of isthmus-dependent atrial flutter. European Heart Journal:
580	Cardiovascular Imaging 2018: October 10. Epub ahead of print.
581	
582	⁵ Dawson DK, Hawlisch K, Prescott G, Roussin I, Di Pietro E, Deac M, Wong J, Frenneaux
583	MP, Pennell DJ, Prasad SK. Prognostic role of CMR in patients presenting with ventricular
584	arrhythmias. Journal of the American College of Cardiology: Cardiovascular Imaging 2013:
585	6; 335-44.
586	
587	⁶ Wijnmaalen AP, van der Geest RJ, van Huls van Taxis CFB, Siebelink H-MJ, Kroft LJM,
588	Bax JJ, Reiber JHC, Schalij MJ, Zeppenfeld K. Head-to-head comparison of contrast-
589	enhanced magnetic resonance imaging and electroanatomical voltage mapping to assess post-

590	infarct scar characteristics in patients with ventricular tachycardias: real-time image
591	integration and reversed registration. European Heart Journal 2011: 32; 104-114.
592	
593	⁷ Desjardins B, Crawford T, Good E, Oral H, Chugh A, Pelosi F et al. Infarct architecture and
594	characteristics on delayed enhanced magnetic resonance imaging and electroanatomic
595	mapping in patients with postinfarction ventricular arrhythmia. Heart Rhythm 2009: 6; 644-
596	51.
597	
598	⁸ Roujol S, Basha TA, Khanna V, Chan RH, Moghari MH, Rayatzadeh H et al. Improved
599	multimodality data fusion of late gadolinium enhancement MRI to left ventricular voltage
600	maps in ventricular tachycardia ablation. IEEE Transactions in Biomedical Engineering
601	2013: 60; 1308-17.
602	
603	⁹ Tschabrunn CM, Roujol S, Nezafat R, Faulkner-Jones B, Buxton AE, Josephson ME, Anter
604	E. A swine model of infarct-related re-entrant ventricular tachycardia: electroanatomic,
605	magnetic resonance and histopathological characterization. Heart Rhythm 2016: 13; 262-73.
606	
607	¹⁰ Mukherjee RK, Roujol S, Chubb H et al. Epicardial electroanatomical mapping,
608	radiofrequency ablation and lesion imaging in the porcine left ventricle under real-time
609	magnetic resonance imaging guidance - an in-vivo feasibility study. Europace 2018; 20: 254-
610	262.
611	
612	¹¹ Tung R, Kim S, Yagishita D, Vaseghi M, Ennis DB, Ouadah S, Ajijola OA, Bradfield JS,
613	Mahapatra S, Finn P, Shivkumar K. Scar voltage threshold determination using ex vivo
614	magnetic resonance imaging integration in a porcine infarct model: influence of

615	interelectrode distances and three-dimensional spatial effects of scar. Heart Rhythm 2016: 13;
616	1993-2002.

618	¹² Brunckhorst CB, Stevenson WG, Soejima K, Maisel WH, Delacretaz E, Friedman PL,
619	Ben-Haim SA. Relationship of slow conduction detected by pace-mapping to ventricular
620	tachycardia re-entry circuit sites after infarction. Journal of the American College of
621	Cardiology 2003: 41; 802-9.
622	
623	¹³ Lopez-Yunta M, Leon DG, Alfonso-Almazan JM, Marina-Breysse M, Quintanilla JG,
624	Sanchez-Gonzalez J et al. Implications of bipolar voltage mapping and magnetic resonance
625	imaging resolution in biventricular scar characterisation after myocardial infarction.
626	Europace 2019: 21; 163-74.
627	
628	¹⁴ Tschabrunn CM, Roujol S, Dorman NC, Nezafat R, Josephson ME, Anter E. High-
629	resolution mapping of ventricular scar: comparison between single and multi-electrode
630	catheters. Circulation: Arrhythmia and Electrophysiology 2016: 9; e003841.
631	
632	¹⁵ Acosta J, Penela D, Andreu D, Cabrera M, Carlosena A, Vassanelli F et al. Multielectrode
633	vs. point-by-point mapping for ventricular tachycardia substrate ablation: a randomized
634	study. Europace 2018; 20: 512-19.
635	
636	¹⁶ Roujol S, Anter E, Josephson ME, Nezafat R. Characterisation of respiratory and cardiac
637	motion from electroanatomical mapping data for improved fusion of MRI to left ventricular
638	electrograms. PLoS One 2013: 8; e78852.
639	

640	¹⁷ Chubb H, Karim R, Roujol S, Nunez-Garcia M, Williams SE, Whitaker J et al. The
641	reproducibility of late gadolinium enhancement cardiovascular magnetic resonance imaging
642	of post-ablation atrial scar: a cross-over study. Journal of Cardiovascular Magnetic
643	Resonance 2018: 20; 21.
644	
645	¹⁸ Jais P, Maury P, Khairy P, Sacher F, Nault I, Komatsu Y et al. Elimination of local
646	abnormal ventricular activities: a new end point for substrate modification in patients with
647	scar-related ventricular tachycardia. Circulation 2012: 125; 2184-96.
648	
649	¹⁹ Acosta J, Andreu D, Penela D, Cabrera M, Carlosena A, Korshunov V et al. Elucidation of
650	hidden slow conduction by double ventricular extrastimuli: a method for further arrhythmic
651	substrate identification in ventricular tachycardia ablation procedures. Europace 2018: 20;
652	337-46.
653	
654	²⁰ Nazarian S, Kolandaivelu A, Zviman MM, Meininger GR, Kato R, Susil RC, Roguin A,
655	Dickfeld TL, Ashikaga H, Calkins H, Berger RD, Bluemke DA, Lardo AC, Halperin HR.
656	Feasibility of real-time magnetic resonance imaging for catheter guidance in
657	electrophysiology studies. Circulation 2008: 118; 223-9.
658	
659	²¹ Oduneye SO, Pop M, Biswas L, Ghate S, Flor R, Ramanan V, Barry J, Celik H, Crystal E,
660	Wright GA. Post-infarction ventricular tachycardia substrate characterisation: a comparison
661	between late enhancement magnetic resonance imaging and voltage mapping using a MR-
662	guided electrophysiology system. IEEE Transactions in Biomedical Engineering 2013: 60;
663	2442-9.
664	

665	²² Oduneye SO, Pop M, Shurrab M, Biswas L, Ramanan V, Barry J, Crystal E, Wright GA.
666	Distribution of abnormal potentials in chronic myocardial infarction using a real-time
667	magnetic resonance guided electrophysiology system. Journal of Cardiovascular Magnetic
668	Resonance 2015: 17;27.
669	
670	²³ Schmidt EJ, Watkins RD, Zviman MM, Guttman MA, Wang W, Halperin HA. A magnetic

resonance imaging-conditional external cardiac defibrillator for resuscitation within the 671

672 magnetic resonance imaging scanner bore. Circulation: Cardiovascular Imaging 2016: 9; 673 e005091.

674

675 ²⁴ Tse ZT, Dumoulin CL, Clifford GD, Schweitzer J, Qin L, Oster J, Jerosch-Herold M,

676 Kwong RY, Michaud G, Stevenson WG, Schmidt EJ. A 1.5T MRI-conditional 12-lead

electrocardiogram for MRI and intra-MR intervention. Magnetic Resonance in Medicine 677 678 2014: 71; 1336-47.

679

²⁵ Elbes D, Magat J, Govari A, Ephrath Y, Vieillot D, Beeckler C, Weerasooriya R, Jais P, 680

Quesson B. Magnetic resonance imaging-compatible circular mapping catheter: an in vivo 681 682 feasibility and safety study. Europace 2017: 19; 458-64.



Figure 1: Set-up of the real-time MR-EP system to enable electrophysiology studies inside a MRI scanner.



Figure 2: Representative depiction of image guidance platform showing 3 orthogonal MRI views demonstrating location of catheter in relation to LV endocardium, 3D segmentation of the left ventricle derived from MRI and scar segmentation from LGE images to guide EAM.



Figure 3: Representative examples of intra-cardiac EGMs obtained using the MR-EP system in a region of normal myocardium (Point 1) and area of scar (Point 2) (A-C). The baseline noise level inside the MRI scanner was in the region of 0.1mV (approximately 10-fold higher than that in the conventional electrophysiology laboratory). Dot-plot showing signal-to-noise ratios obtained for intra-cardiac EGMs in normal myocardium and LGE-derived scar regions from 7 animals; *p<0.0001(D).



Figure 4: Representative examples of segmented LGE scar, voltage and S-QRS maps obtained using real-time MR-EP system in 2 animals (Panel A and B)



Figure 5: ROC curves (A and C) for prediction of LGE regions using voltage mapping and S-QRS. Frequency histograms (B and D) displaying the true positive, false positive, true negative and false negative counts of voltage mapping and S-QRS measurements using the real-time MR-EP system to predict MRI-derived scar. Sensitivity, specificity, PPV and NPV of measurements using the system using different normal voltage cut-offs and S-QRS times (E-H)



Figure 6: Sorenson-Dice similarity co-efficient between MR-derived scar shells (far left panels) and endocardial voltage maps with varying normal voltage thresholds (right panels) in 2 representative animals.



Figure 7: Dice similarity co-efficients (DSC) between MR-derived scar shells and endocardial voltage maps acquired using MR-EP system following application of normal cut-off thresholds of 0.5-3.5mV. DSC is shown for overall similarity, similarity across scar nodes and normal myocardium nodes.