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Cardiac MRI Endpoints in Myocardial Infarction Experimental and Clinical Trials

JACC Scientific Expert Panel

Borja Ibanez, MD, PnD,a,b,c Anthony H. Aletras, PnD,d,e Andrew E. Arai, MD, f Hakan Arheden, MD,g Jeroen Bax, MD, PnD, h Colin Berry, MD, PnD, i Chiara Bucciarelli-Ducci, MD, PnD, j Pierre Croisille, MD, PnD, k Erica Dall’Armellina, MD, PnD, l Rohan Dharmakumar, PnD, m Ingo Eitel, MD, n Rodrigo Fernández-Jiménez, MD, PnD, o,p Matthias G. Friedrich, MD, q,r,s David García-Dorado, MD, t,u Sebastian Kozerke, PHD, v Christopher M. Kramer, MD, w Michael Salerno, MD, PnD, MS, x Javier Sánchez-González, PnD, y Javier Sanz, MD, z Valentin Fuster, MD, PnD, ab

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ABSTRACT

After a reperfused myocardial infarction (MI), dynamic tissue changes occur (edema, inflammation, microvascular obstruction, hemorrhage, cardiomyocyte necrosis, and ultimately replacement by fibrosis). The extension and magnitude of these changes contribute to long-term prognosis after MI. Cardiac magnetic resonance (CMR) is the gold-standard technique for noninvasive myocardial tissue characterization. CMR is also the preferred methodology for the identification of potential benefits associated with new cardioprotective strategies both in experimental and clinical trials. However, there is a wide heterogeneity in CMR methodologies used in experimental and clinical trials, including time of post-MI scan, acquisition protocols, and, more importantly, selection of endpoints. There is a need for standardization of these methodologies to improve the translation into a real clinical benefit. The main objective of this scientific expert panel consensus document is to provide recommendations for CMR endpoint selection in experimental and clinical trials based on pathophysiology and its association with hard outcomes. (J Am Coll Cardiol 2019;74:238–56)

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objectives of this 2-day summit were to appraise prior and new lines of evidence of CMR-based depiction of myocardial I/R injury, to review available data on the diagnostic and prognostic value of CMR in the post-MI setting, and to provide recommendations for standardization of imaging protocols for use in clinical trials and experimental scenarios. This Expert Panel consensus document summarizes the final conclusions and recommendations agreed in this meeting. Philips sponsored the meeting (travel of experts and meals, no economic compensation to any person involved in this meeting) but did not play any role in the appointment of experts, topics or conclusions. The process of appointing the expert panel and the methodology followed for the identification of topics and for deciding statements/recommendations is presented in the Online Appendix. Although there are other relevant recent consensus documents from scientific societies dealing with methodologies and recommendations on CMR outcome measures (2–5), the present consensus is only focused on MI (I/R). The document addresses 2 main areas: 1) pathophysiology of myocardial I/R injury as revealed by CMR; and 2) proposed endpoints for use in clinical and research studies in the post-MI setting. It is important to highlight that although some pathophysiology and imaging principles apply to both STEMI and non-STEMI, this paper focuses on STEMI.

THE INFARCTED MYOCARDIUM IS NOT STATIC: EVOLVING CHANGES AFTER I/R

A section dedicated to the relevance of animal models to human pathophysiology and how these contribute to our knowledge about I/R can be found in the Online Appendix.

PROGRESSION OF NECROSIS AND ITS BORDERS. In the late 1970s, Reimer and Jennings (6) experimentally demonstrated in dogs that necrosis ensues as a “wave front” from the subendocardium to the epicardium, following a transmural gradient of collateral flow within the hypoperfused (ischemic) area. Although some investigators initially suggested the existence of a wide lateral “border zone” of intermediate-level perfusion, later studies showed this was mostly a partial volume artifact due to the limited resolution of the techniques used, and indeed, with progressively higher levels of spatial resolution, investigators have concluded that there is no zone of intermediate perfusion (or injury) at the lateral border of the area at risk (AAR) in dogs (7,8). Despite the prevailing concept based on pathology studies that there is no lateral progression of necrosis, there are some studies suggesting the opposite in some cases (9–11). Given that most of these studies used CMR and not pathology, these data should be interpreted with caution.

At the infarct border, histological sections can show apparent isolated islands of surviving myocardium within areas of necrotic tissue, as well as isolated islands of necrosis within areas of viable tissue (12). In the absence of collaterals, the infarct borders are sharp at a cellular level but may appear to have fuzzy borders when assessed by methods with low resolution relative to cardiomyocyte size due to partial volume errors (13).

DYNAMIC CHANGES OF INFARCT SIZE AFTER I/R. When assessing the post-I/R myocardium by CMR, the imaging features of the “infarcted myocardium” can experience important longitudinal changes. Immediately after reperfusion, the area that will become irreversibly injured increases in size due to reperfusion-related edema. This “swelling” is rapidly reduced, resulting in a decrease in the size of the irreversibly injured myocardium. In the following days, dead cardiomyocytes and debris are progressively removed from the post-I/R region, while there is an intense transit of inflammatory cells and a
progressive deposition of collagen. These changes result in a further "shrinkage" of the infarcted region (Figure 1). Depending on the extent of irreversible injury and its spatial distribution (e.g., subendocardial, transmural, or patchy), these changes can be of different magnitudes. This concept is important, because the imaging biomarker "infarct size" (as % of LV) can vary greatly at different time points after MI (14–16). Studies aiming to quantify MI size should be consistent in the choice of the imaging time point to reduce variability and bias in outcome measure.

**PATHOPHYSIOLOGY AND SPATIAL DISTRIBUTION OF POST-I/R EDEMA.** In the healthy heart, myocardial water content is mostly intracellular, with a very small interstitial component (17). Upon MI, myocardial edema occurs initially as cardiomyocyte swelling.

**TABLE 1 Consensus Related to Myocardial I/R Concepts and Animal Models**

| Concept/Animal Models |
|-----------------------|
| Small animal models (mice, rats) are useful for specific mechanistic evaluations but do not resemble human pathophysiology. It is not recommended to translate results from these models into humans without first testing in large animals (126). |
| Pig and dog I/R models have similarities and differences with humans but are the closest to human pathophysiology and may be considered complimentary* (primates are not considered for ethical reasons) (126). |
| The post-I/R region (infarcted and salvaged) is not static and exhibits important temporal changes in size and composition (14). |
| There is greater edema formation in infarcted tissue than in salvaged myocardium (22,70). |
| After acute MI, the infarct border can be highly irregular with islands and peninsulas of necrosis but with a sharp boundary and no "intermediate zone" at a cellular level (12). "Fuzzy borders" at the edge of an acute infarct can arise from partial volume effect due to limited spatial resolution of imaging (13). |
| In experimental models, edema after I/R is bimodal (70,80,127). There is an initial reperfusion-related wave of edema that significantly attenuates hours after reperfusion. A second healing-related wave of edema initiates days after reperfusion and lasts longer (15,50). |
| In humans, the dynamics of post-I/R edema is less well established. At least in some patients it might exhibit a 2-wave pattern (45,81–83). Edema appears to be more stable in the time window between days 3 and 7 post-MI (15,45,82,83,90). |
| MVO precedes IMH in reperfused infarctions (15,128). |

*Controversial topic (majority but not all panelists in agreement).*

IMH = intramyocardial hemorrhage; I/R = ischemia/reperfusion; MI = myocardial infarction; MVO = microvascular obstruction.
during the early stages of ischemia (18). Edema is then abruptly exacerbated on restoration of blood flow to the ischemic region. This reperfusion-associated increase in edema is caused by increased interstitial (extracellular) edema secondary to rupture of myocytes, reactive hyperemia, and leakage from damaged capillaries (19–21). Myocardial edema observed days or weeks after reperfusion is mostly extracellular, given that energetic recovery is relatively rapid in surviving cardiomyocytes, and related to tissue healing (17).

In relation to the spatial extent of post-MI edema, classic pathology studies have demonstrated larger amounts of edema in the actual infarcted myocardium than in the “reversible” injured tissue (i.e., salvaged myocardium) (18,22). These classical studies showed that infarcted myocardium has an increase in water content of ≤90%, while salvaged myocardium only has an increase of water content of ≤9% for <24 h. As discussed in more detail in subsequent sections, the dynamics of post-MI edema is highly complex.

**PATHOPHYSIOLOGY OF MICROVASCULAR INJURY: MICROVASCULAR OBSTRUCTION AND INTRAMYOCARDIAL HEMORRHAGE.** As the ischemia progresses capillary permeability increases, and upon reperfusion, the increased interstitial pressure overwhelms the intravascular pressure and can lead to microvascular obstruction (MVO) by external compression (1). When reperfusion is delayed, the endothelial walls are compromised at reperfusion by the sudden appearance of a positive pressure in the microvasculature resulting in microvascular destruction causing intramyocardial hemorrhage (IMH). Accordingly, MVO appears to precede IMH in reperfused infarctions, which implies that hemorrhagic infarctions will always show evidence of MVO. MVO has been shown to evolve over time and stabilize by 48 to 72 h (23).

Given that microvascular damage is associated with late reperfusion, it is not surprising that ischemic time is associated with larger MVO and IMH areas. In this regard, a chronic marker of microvascular injury is persistent iron deposits, which have
been shown to occur in large animal models and in humans (24–26). These chronic iron deposits have been associated with significant prolonged proinflammatory burden, poor LV remodeling, and major adverse cardiovascular events. Thus, markers of severe forms of tissue injury, particularly MVO and IMH, appear to be key targets for post-infarction therapy (27). See Table 1 for consensus related to I/R concepts and animal models.

**POST-MYOCARDIAL INFARCTION TISSUE CHARACTERIZATION BY CMR**

The aim of this section is to provide an overview of CMR methods for imaging the processes occurring in the post-I/R myocardium. For a technical description of the methods, the reader is referred to contemporary Society for Cardiovascular Magnetic Resonance consensus documents (2–5).

**CMR IMAGING OF INFARCT SIZE.** Late gadolinium enhancement. Ti-weighted (T1W) late gadolinium enhancement (LGE) has clearly been established as the best in vivo surrogate for infarct size. Gadolinium-chelates are extracellular contrast agents. Following intravenous administration of gadolinium, the contrast agent perfuses locally through myocardial capillaries and diffuses within the interstitial compartment, reaching a dynamic equilibrium allowing quantification of the extracellular volume (ECV) (28). In the early post-infarcted tissue, the rupture of cardiac cells increases the interstitial volume, which reduces the washout rate, and results in an increase in contrast concentration compared with healthy tissue; gadolinium persists within the infarct zone for a longer time than in unaffected tissue, leading to a regional hyperintense (bright) area on T1W imaging (Figure 2). At later post-MI stages, dead cardiomyocytes are replaced by extracellular matrix resulting in chronic increased ECV, also resulting in increased volume of distribution of gadolinium with similar imaging effects. Contrast-enhanced LGE typically involves breath-hold inversion-recovery imaging 10 to 20 min after intravenous administration of a gadolinium-based contrast agent. 3-Dimensional (3D) navigator-gated LGE acquisitions are now feasible expanding volumetric coverage of myocardial tissue (29). 3D single breath-hold acquisitions have the advantage of a better delineation of complex lesions, with greater signal intensity and contrast-to-noise ratio compared with 2-dimensional approaches (30). Expanded information about LGE imaging principles can be found in the Online Appendix.

**Effect of infarct size dynamics and partial volume effects on LGE interpretation.** The size of apparently “dead” myocardium may evolve over time, and this is an important consideration when interpreting LGE images (14–16). The shrinkage in LGE size between a scan performed in the early days after MI and a repeat scan performed days to weeks later may be misconstrued as evidence that LGE represents both viable and nonviable myocardium acutely, whereas LGE represents only nonviable myocardium chronically. Given the rapidly dynamic tissue changes happening acutely, the validity of a direct comparison between LGE size and histological infarct size is arguable if quantification with each method is not performed at the same time/day.

Infarct involution during healing can also interact with partial volume effects to accentuate changes that might occur in LGE size from the acute to chronic setting. Voxels containing islands or peninsulas of necrosis at the infarct border are often partially “bright” (or “grey”) on LGE imaging and may be above the arbitrary threshold used to define abnormal
myocardium. But, at a later time point, shrinkage of these peninsulas or islands leads to a changed partial volume effect, and voxels at the infarct border may no longer be considered hyperenhanced since they are now below the cutoff threshold (31) (Figure 3).

For certain types of injuries, the combination of infarct shrinkage and partial volume effect can lead to LGE findings that are particularly prone to misinterpretation. If there is patchy necrosis, hyperenhancement in the acute setting may resolve and be absent in the chronic setting (Figure 2).

Given the dynamics of acutely infarcted myocardium, there are several implications to the design and interpretation of CMR studies evaluating infarct size. Differences in size of injured myocardium as assessed by the same or different imaging techniques on separate days should not be considered to reflect different underlying pathophysiological components without accounting for the dynamic changes in size. Similarly, size-based findings on CMR should not be compared to a pathology reference standard unless both were performed on the same day.

Pre- and post-contrast T1 mapping for assessment of acute infarct size. ECV mapping is determined by obtaining repeated T1 maps before (native T1 map) and after contrast media administration (32). ECV agrees well with final infarct size revealed by LGE (32). Despite the fact that native T1 and ECV mapping might overcome some of the limitations of LGE imaging, it is subjected to the same considerations regarding infarct size dynamics. In addition, the spatial resolution of ECV mapping is inherently less than that of LGE imaging.

A section dedicated to post-processing of LGE and edema images is presented in the Online Appendix.

CMR IMAGING OF EDEMA. The identification of the myocardial edematous regions has become part of clinical and research CMR studies for different cardiac conditions (mainly myocarditis and post-MI). For several years, the extent of post-MI edema (as identified by different CMR methodologies) has been used as a surrogate for AAR and thus utilized in a multitude of experimental and clinical studies aiming at quantification of myocardial salvage. The concept of post-MI edema extension as equivalent to AAR has recently been challenged, and there are many uncertainties about its accuracy. Conversely, the degree (and extent) of post-MI edema is increasingly acknowledged as a marker of injury itself.

CMR as a tool to detect myocardial edema. CMR offers the possibility of identifying areas of increased myocardial free water content or edema (3). The differentiation between intracellular and extracellular edema is technically challenging; however, CMR better reflects an increase in the extracellular compartment (33). As the extracellular compartment is the one showing the largest change in water content after MI, CMR closely correlates with changes in total myocardial water content (34,35). Nevertheless, there might be other factors affecting the edema-related post-MI CMR signal, such as changes in water fractions (protein-bound vs. free) (36) or the paramagnetic effect of hemoglobin denaturation in the presence of tissue hemorrhage (37). Here, a brief description of different CMR sequences for edema imaging is presented. More extensive information can be found in the Online Appendix.

T2-weighted pulse sequences. The transverse relaxation time (T2) directly reflects tissue composition and hydration. Hyperintense zones on T2-weighted (T2W) CMR images are indicative of myocardial edema. The post-MI edema formation, as assessed both ex vivo and in vivo (34,38), has been shown to correlate with acute myocardial injury (39–45). By allowing the detection of edema post-acute ischemic injury (46), T2W-CMR might help its discrimination from chronic MI (41,47).

Dark-blood T2W-CMR methods are widely used for clinical and research purposes (42–45,48–52). More recently, bright blood T2W-CMR techniques have emerged as potential alternatives to dark blood T2W-CMR (53–55).

T2 mapping. Parametric T2-mapping techniques provide voxel-based tissue characterization on a continuous quantitative scale. By displaying T2 values in milliseconds on a fixed scale, subjective signal intensity-based ascertainment is avoided. T2 mapping has been validated against pathology for myocardial water content quantification in experimental MI models (35,56). A comparative study using repeatedly 4 different edema sequences showed higher intraobserver and interobserver agreement and greater test-retest reproducibility in the quantification of edema using T2 mapping (57).

T1 mapping. When the proton is contained in a large molecule, such as fat, energy transfer is very efficient, resulting in short T1. On the contrary, for small molecules, such as water, this effect is less efficient, resulting in longer T1. Myocardial injury due to MI or inflammation increases water content, lengthening local T1 relaxation times (3). The extent of edema delineated by T1 maps agrees well with other measures (58–60).

Retrospective AAR size estimation by CMR: an evolving controversy. The “AAR” concept. The acutely ischemic myocardium downstream of an
occluded coronary artery defines the myocardium at risk of death, the so-called AAR. The portion of the AAR that survives, if any, represents myocardium that has been salvaged (eventually by a given intervention).

Although infarct size provides an approximate (inverse) estimate of myocardial salvage, the extent of the AAR is highly variable and dependent on the infarct-related artery, coronary distribution, and location of the culprit lesion. Thus, a better measure of therapeutic efficacy would be the proportion of the AAR that is infarcted rather than absolute infarct size. The rationale is that for 2 equally sized infarcts, the one representing a larger proportion of the AAR corresponds to a greater amount of salvaged tissue (61) (Figure 4). Thus, the quantification of infarct size relative to AAR theoretically overcomes the limitation of interindividual AAR extent variability and, thus, might allow an accurate estimation of the effectiveness of protective therapies while reducing the necessary sample size. This methodology, despite being conceptually correct, is dependent on the
accuracy of the technique(s) used for AAR and infarct size quantifications.

**Dynamic changes in the AAR of the Acutely Infarcted Myocardium.** The post-I/R region (the AAR) is usually composed of separate myocardial compartments that are biologically diverse. For instance, in the setting of a subendocardial infarction, the necrotic tissue in the subendocardial zone of the AAR has a different biological milieu than the viable tissue in the subepicardial zone. Besides a large difference in electrolytes, metabolites, inflammatory cells, and many other biochemical and biophysical properties, the underlying structure by light and electron microscopy are wholly dissimilar (18,19,22,62,63). This raises reservations regarding the use of such characteristics to provide an accurate depiction of the AAR (61).

Another important concept to take into consideration in this regard is that the tissue composition of the post-I/R region (the AAR) is dynamic in composition. Within the infarcted component, there is a clear time course of histopathological changes as necrotic myocardium heals into collagenous scar (50). Many features, such as acute inflammation, may not be stable even within the first week. This has consequences for imaging the AAR if the imaging method is dependent on a highly dynamic attribute of MI.

**Edema-based CMR methodologies for retrospective AAR size estimate.** Landmark experimental studies suggested the possibility of ex vivo (64) and in vivo (38) retrospective (i.e., after I/R) AAR quantification by noncontrast T2W-CMR (edema). The first clinical study on this topic showed that after MI, areas of increased T2 signal extended beyond that of areas of LGE (39), showing that T2W-CMR of myocardial edema could detect reversibly injured myocardium in STEMI patients. After these pioneer works, several experimental and clinical studies have studied the relationship between the extent of edema-CMR abnormalities (T2W, T2 mapping, T1 mapping) and different AAR standards showing positive associations (59,60,65-72). There were also initial experimental studies concluding against the validity of edema CMR imaging for retrospective AAR quantification (20,44).

One limitation to most of these studies is that CMR imaging was performed at a single time point after reperfusion under the assumption that edema was stable for several days. The second assumption of these studies was that edema was not affected by cardioprotective strategies and/or experimental conditions (e.g., duration of ischemia). Recent experimental and clinical evidences have challenged both assumptions, and thus, there is uncertainty regarding the concept that edema equates AAR across different experimental/clinical scenarios.

Contrast-based CMR methodologies have also been proposed for retrospective AAR quantification: early gadolinium enhancement (73), ECV imaging (70), and contrast-enhanced steady-state free precession (10,74-76) (Online Appendix). The mechanistic explanation for the possible retrospective AAR estimation by these contrast-based methods is not clear. The advantage of contrast enhanced steady-state free precession is that it does not require any additional sequence.

CMR methods for retrospective AAR quantification not relying on edema have been proposed: the endocardial extent of infarction on LGE images (termed infarct-endocardial surface length) is based on the wave front progression of necrosis. The relationship between this method and different standards have been tested with positive results (77,78). However, other studies are questioning its accuracy in STEMI patients with minimal or no LGE (11,67,68).

**Dynamics of post-MI edema: the classic concept of “stable” myocardial edematous reaction revisited.** Post-MI edema was initially believed to be stable during the days following an acute MI (14,66). Recent studies have provided new insights. In the experimental setting, it has been demonstrated that the edematous reaction during the first week after MI is not stable, and follows a bimodal pattern (79,80). The 2 waves of edema are secondary to different pathophysiological phenomena (Figure 1), and are amenable to being modified by interventions (15,21,50).

In a longitudinal clinical study of 16 STEMI patients, including CMR within the hyperacute post-reperfusion period (i.e., 2 h after reperfusion), the post-MI bimodal edema pattern was observed in a significant proportion of patients (45). Infarcted (either with or without IMH) and, to a lesser extent, “reversibly” injured myocardium displayed a similar 2-wave time course (45), as shown in the experimental setting (15,70). In another longitudinal study of 30 STEMI patients, edema was shown to be stable across different time points ≥10 days in all patients except those with IMH (81). Given the dynamic changes in the post-MI myocardium early after reperfusion, the discrepancies between these studies might be explained by the different post-reperfusion timings of the CMR scans between studies. More recently, a third clinical study showed that intensity of post-STEMI edema according to T1 mapping relaxation times was bimodal, while the extent of T1
TABLE 2  Consensus Related to CMR Methodologies for Myocardial Tissue Characterization

Conceptually, the AAR is the myocardium downstream to an occluded coronary artery that becomes ischemic (1). Infarct size and AAR should be ideally measured at the same time point. If measured at different time points, salvage quantification might show implausible results (e.g., negative values) (14-16,90).

LGE grey zones should not be interpreted as dead, since they likely represent a mixture of bright (dead) and dark (alive) myocardium. Grey myocardium can be due to partial volume effect (15,31).

Edema development should be interpreted as a manifestation of myocardial I/R injury and should not be considered as a reliable marker of AAR* (15,44,89).

T1W is the recommended methodology for LGE imaging after MI (2,4).

T2-mapping is the recommended methodology for edema imaging after MI (3,35).

T1W LGE is the recommended methodology for MVO imaging (hypointense areas within the LGE area) (95).

T2* mapping is the recommended methodology for IMH imaging after MI (3).

*Controversial topic (majority but not all panelists in agreement).

AAR = area at risk; LGE = late gadolinium enhancement; other abbreviations as in Table 1.

abnormalities was not (82). Dynamic tissue water changes (T2 values) were also suggested in a recent longitudinal clinical study (83), but not in other retrospective studies (76,84,85).

In summary, there is robust evidence for the bimodal edema pattern at the experimental level. In patients, some (not all) studies have also demonstrated a bimodal edema reaction within the first week of MI. **Effect of protective therapies on edema: implications for edema-based AAR quantification.** Thuny et al. (86) were the first group to study the effect of a cardioprotective strategy (post-conditioning) on the extension of edema, evaluated by T2W abnormalities. A total of 50 STEMI patients were randomized to the cardioprotective maneuver post-conditioning or control. The extent of edema (T2W) was significantly smaller in the post-conditioning group. In other studies where post-conditioning was not associated with protection, the effect of this maneuver on edema was not observed (15,87). In a subsequent clinical study, White et al. (88) evaluated the effect of remote ischemic conditioning (RIC) on infarct size (LGE extension) and edema development (T2 mapping) in 83 STEMI patients. The extent of edema on T2 mapping was significantly smaller in the group undergoing RIC. In addition, mean T2 relaxation times in the ischemic area were significantly shorter in the group of patients undergoing RIC compared with control.

In the pig model of I/R, Fernandez-Jimenez et al. (15) evaluated the effect of the cardioprotective maneuver local ischemic preconditioning (IPC) on edema formation. The extension of edema was significantly smaller in pigs undergoing IPC than in controls. T2 relaxation times at the ischemic area were significantly shorter in IPC pigs (15). Some degree of edema is still present days after MI in the absence of LGE (15), suggesting that edema might be a marker of injury. In a different study undertaken in a mouse model, Dongworth et al. (65) investigated the effect of IPC on edema development assessed by T2 mapping and found that IPC reduced the extent of edema (T2 mapping).

In agreement with these experimental data, very recently Bulluck et al. (89) reported a meta-analysis of randomized clinical trials testing the infarct-limiting effect of different interventions using CMR within the first week after STEMI. A total of 5 trials reported that the active intervention resulted in smaller infarctions, and in these, the extension of edema (T2W abnormalities) was also significantly reduced. Conversely, in trials where the active intervention was not associated with smaller infarctions than controls, the extension of edema was not different either (89).

Overall, there is compelling experimental and clinical data showing that cardioprotective therapies potent enough to reduce irreversible injury (i.e., infarct size), also limit the size and degree of edema. Duration of ischemia also affects edema formation following acute MI. Edema development, therefore, should be seen as a manifestation of myocardial I/R injury and not a simple marker of AAR.

The assumption that the extent of edema after myocardial infarction accurately represents AAR in all experimental/clinical scenarios is under intense debate, and thus, edema-based methodologies for AAR assessment in testing novel cardioprotective therapies cannot be recommended. The panel wants to make clear that this conclusion and recommendation does not mean that seminal studies showing correlations between CMR-based edema extension and standard-based AAR were not correct. These studies were performed with only 1 experimental condition (fixed ischemia duration without testing cardioprotective strategies), and at 1 single time point after I/R. This recommendation should not be interpreted as the last nail in the coffin of the field of using edema as surrogate for AAR. Further ad hoc research must be undertaken to better understand the interplay between injury and edema formation/dynamics. **Salvage index concept and its quantification.** The most commonly used and robust CMR measurement for assessing the efficacy of novel cardioprotective therapies in STEMI patients is an absolute reduction in
infarct size. However, given the variability in the size of the AAR, the number of individuals required in clinical cardioprotection studies to demonstrate a reduction in absolute MI size can be quite high (approximately 200 to 300 STEMI patients in total depending on the patient population and effect size) (90). Normalization of infarct size by the AAR theoretically reduces variability of outcome measure and thus considerably reduces sample size. Thus, myocardial salvage index (MSI) is defined as the proportion of the AAR that remains viable after MI.

In the experimental setting, the gold-standard method for delineating the AAR in animal models of I/R is to quantify the perfusion defect following injection of either fluorescent microspheres or dyes (such as Evans blue). Alternatively, an arterial enhanced multidetector computed tomography (MDCT) performed during the index coronary occlusion is a valid technique to delineate the theoretical AAR in vivo (15,20,72). In the clinical setting, the gold-standard method for quantifying the perfusion defect in STEMI patients is by cardiac single-photon emission computed tomography imaging of the perfusion defect within hours after reperfusion following the peripheral injection of a radioisotope (e.g., technetium) prior to opening of the infarct-related artery (66).

In the experimental setting, MSI can be quantified pathologically (tetrazolium chloride and microspheres/blue dyes for infarct and AAR quantifications, respectively), but this is not possible in the clinical setting. To overcome these limitations,
CMR has been used to quantify both infarct size (LGE) and AAR in a single imaging session. Based on the assumption that T2-based CMR sequences were identifying the AAR and that LGE is a valid surrogate for infarct size, CMR-based MSI has become a popular outcome measure in experimental and clinical trials. The use of CMR-based MSI as the primary endpoint in trials has been proposed as a methodology to allow a reduction in sample size by one-half (91). Studies have proposed that CMR-MSI calculated by a single CMR in the first week following STEMI can predict long-term clinical outcomes (90), suggesting that it is a viable surrogate endpoint for assessing cardioprotection. Based on the recent evidences highlighting that edema is not stable after MI, and that cardioprotective interventions may affect the development of edema, edema-based MSI should not be used as an endpoint in experimental or clinical trials. Similarly, endocardial surface length LGE methodologies for AAR assessment in testing novel cardioprotective therapies should be used with caution, since they are not reliable in cases of very limited LGE, and are not recommended as a primary or secondary outcome measure.

**CMR IMAGING OF MICROVASCULAR INJURY: MVO AND INTRAMYOCARDIAL HEMORRHAGE. Microvascular obstruction.** In experimental studies, the intracoronary administration of dye (Thioflavin-S) may result in an excellent delineation of MVO in the excised heart (92), but this is not feasible in the clinical setting. On CMR, MVO is revealed as a hypointense core within the hyperintense infarct zone within the area of LGE (93–95). The imaging feature is explained by failure of the contrast agent to penetrate within the infarct core.

Native T1 mapping may allow the detection of areas with short T1 values surrounded by prolonged T1 values in the infarcted regions. Although initially thought to be reflective of MVO, recent evidence has led to a definition of a new biomarker of “infarct core” (96). The mechanisms underpinning the shortening of T1 values are not fully understood.

**Intramyocardial hemorrhage.** Deoxyhemoglobin has paramagnetic effects that enable myocardial iron accumulation to be detected using T2W and T2*W CMR (97,98). IMH is visually defined as a hypointense area in the center of a hyperintense zone in edema sequences. Nevertheless, T2*-based imaging is the most sensitive method for imaging hemorrhage as it specifically images IMH based on reduced T2* relaxation times. Despite this, given that spatial resolution is relatively low; it is also sensitive to off-resonance artifacts, particularly in the lateral wall; and it is influenced by motion and blood flow, T2* CMR image quality can be suboptimal. Two different cut-offs are used to identify zones of hemorrhage (mean-2SD and T2* <20 ms). Whether both of the cut-offs yield the same or different hemorrhage volume and whether this depends on the imaging field strength remains to be investigated. An alternative to T2* imaging that has been extensively investigated is T2 imaging, as T2 images can be sensitized to hemorrhage. Although T2 images often provide far better image quality than T2* images, their sensitivity in the acute phase to hemorrhage is compromised by the presence of

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**TABLE 3 Consensus Related to CMR Endpoints in Myocardial Infarction Experimental and Clinical Studies**

It is recommended that the index CMR scan be performed 5 ± 2 days after reinfusion. At this time point, several key parameters have been shown to be relatively stable.

LGE extent, expressed both in absolute (grams) and relative (percentage of LV mass) terms, is the recommended CMR primary endpoint (106,107). In experimental studies, if LGE extent is expressed as a ratio to AAR, the latter should not rely on edema and ideally both AAR and LGE have to be obtained at the same day post-MI (e.g., perfusion imaging during coronary occlusion by MDCT) and LGE by CMR (15,89,90).

LVEF (%) and MVO (areas of hyperenhancement within LGE, in grams or as % of LV) are recommended as main secondary endpoints (105,108).

Edema extent (T2 mapping) may be used as secondary endpoint to assess the effect of cardioprotective therapies (89).

LV volumes (ml; ml/m2), RVEF (%), RV LGE (presence), and IMH (% of LV) can be included as secondary endpoints (111–114).

Myocardial mapping indexes (T1 [ms]; T2 [ms]; ECV [percentage of LV mass]), and strain (feature tracking) may be considered as exploratory endpoints (15,65,88,117–119).

The use of edema-sensitive CMR sequences as a surrogate for AAR (as well as edema-based MSI) is not recommended because edema is not stable in the days following MI, and is potentially affected by duration of ischemia, and by cardioprotective strategies* (15,45,89).

Due to its easy implementation, ESL-based salvage index may be used as an exploratory outcome but not in cases with small amounts of LGE or aborted infarction, i.e., when a high degree of cardioprotection is expected* (11,67,68,77,78).

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For LGE post-processing, automated techniques using signal-intensity thresholding, manual delineation, and visual scoring (the 2 latter at intermediate signal-intensities) have similar good accuracy and reproducibility (129). For delineation of edema, T2 mapping (automated >2SD thresholding with manual correction) showed greater intraobserver and interobserver agreement (57). For MVO quantification, the most widely used is the manual delineation of dark areas within LGE region. Extended information is provided in the Online Appendix. *Controversial topic (majority but not all panelists in agreement).
edema in the acute phase. Because $T_1$ is sensitive to the degradation products of hemorrhage, some studies have also investigated the capacity to image hemorrhage with $T_1$ CMR. These studies have shown that the opposite effects hemorrhage and edema impose of $T_1$, the sensitivity for detecting hemorrhage based on $T_1$ after MI is often compromised (99). See Table 2 for consensus related to CMR methodologies for tissue characterization.

**CMR ENDPOINTS IN CLINICAL TRIALS AND EXPERIMENTAL STUDIES**

Large animal experimentation is necessary before embarking in clinical trials testing new cardioprotective strategies. Given that all imaging methodologies used in patients can be used in large animal studies, to improve transferability of experimental results, CMR endpoints in the latter should be the same as in clinics. For this reason, all statements and conclusions in this section apply both to clinical trials and large animal experimental studies. Additional information about experimental study recommendations is provided in the Online Appendix.

Besides the tissue characterization abilities, the high reproducibility of CMR makes it the best imaging modality for MI clinical trials. Interscan reproducibility for the quantification of LV volumes, LV mass, and LVEF are superior to that of echocardiography, enabling 55% to 93% reductions in sample size to detect changes in these variables with comparable power (100). Similarly, LGE is highly reproducible in both the acute and chronic MI setting (mean bias <1% of LV mass) (101), which allows reductions of approximately 42% of sample size in comparison with single-photon emission computed tomography (102).

To incorporate new interventions into clinical practice, it must be demonstrated that these improve clinical outcomes, particularly “hard” events, such as death or relevant morbidity (HF, use of resources, and so on). However, such trials typically require very large sample sizes and/or follow-up, particularly in the current era of continuously improving post-MI prognosis, and frequently result in prohibitive costs and/or logistic hurdles. Therefore, surrogate endpoints with proven links to hard outcomes are often used as an alternative in early (pilot) clinical trials. A number of CMR-based endpoints derived from dynamic (“functional”) or stationary (“morphologic”) imaging have been proposed as valid surrogates. Although most studies have examined the prognostic value of individual CMR measures, some have combined several into scores (103,104).

In 2014, El Aidi et al. (105) performed a systematic review of 27 studies evaluating the ability of CMR to predict outcomes when performed within 2 weeks of reperfused MI. According to an arbitrary definition, no single CMR measure demonstrated independent prognostic value for the prediction of hard endpoints (death, MI, or transplantation) with the exception of LVEF, which was the only endpoint independently associated with major adverse events. Consistent prognostic value, although insufficient to qualify as “independent,” was additionally noted for LGE size and MVO.

Reduction of MI size is the ultimate goal of cardioprotection in the acute phase. A subsequent pooled analysis of 10 randomized clinical trials evaluated the prognostic significance of imaging-based MI size measured at a median of 4 days post-reperfusion (106). This study demonstrated clear progressive rises in death or HF hospitalization with increasing MI size, even after adjusting for relevant factors. Additional analyses indicated an association between therapy-related reductions in MI size with decline in HF hospitalizations, but not mortality (107).

A more recent analysis of the same randomized trials identified MVO (measured at a median of 3 days post-reperfusion and present in 57% of patients) as an additional independent predictor of mortality after accounting for clinical characteristics and LGE size (108).

In their systematic review, El Aidi et al. (105) found insufficient evidence for any incremental value of other CMR-derived parameters such as segmental wall motion or perfusion abnormalities (at rest and/or stress), edema, or IMH. Although newer reports are available since 2014, particularly regarding IMH, edema, or MSI (103,109), overall evidence for an independent prognostic value for these indexes is far less robust and consistent than for LGE, LVEF, and MVO. Of note, CMR-based MSI on CMR performed 1 to 4 days after STEMI was proposed to entail long-term prognostic value in a cohort of 202 patients (110). However, as discussed in this consensus document, the use of post-MI edema extent as a surrogate for AAR is troublesome (15). Similarly, few small studies suggested potential prognostic implications of alternative measures such as LV volumes (111,112), right ventricular EF (113), right ventricular LGE (114), and more recently, the evaluation of LV strain by feature tracking CMR (115-117).

The extent of edema after reperfusion (as evaluated by different T2 sequences [15]), as well as the
signal intensity (T2W) (86) and T2 relaxation times (15,65,88), can be affected by strong cardioprotective interventions. Therefore, edema development might be used as a secondary endpoint as a marker of injury (not as a surrogate for AAR).

Newer parametric (mapping) techniques evaluating changes in T1, T2, and/or ECV in both infarcted and remote myocardium hold substantial promise for more objective, operator-independent tissue characterization. Although these variables have generally been linked to adverse LV remodeling, there is presently a paucity of data regarding their associations with clinical events, except for some early evidence for native T1 values of remote myocardium (118,119).

In terms of the best post-MI timing for CMR performance in clinical trials, some key parameters holding robust association with hard endpoints (LGE, MVO) change over time (45,90). To be able to quantify different CMR parameters included as endpoints in the same imaging session, and based on available data, the ideal post-MI timing for the evaluation of CMR endpoints appears to be between days 3 and 7 (45,90). LGE and MVO show relative stability and proven prognostic value in this time window.

The Central Illustration shows the hierarchy of recommended CMR outcomes in clinical trials and experimental studies. See Table 3 for consensus related to CMR endpoints in myocardial infarction experimental and clinical studies.

**FUTURE DIRECTIONS**

In the Online Appendix, there is a section presenting several evolving CMR methodologies that may offer insights into tissue characterization and LV function following STEMI that are either already being investigated or have the potential to be applied to STEMI patients.

CMR has become an indispensable technique for post-MI tissue characterization. However, further work is required to make CMR simpler, faster and more standardized to realize the ultimate goal of individualized patient management post-MI, tailored according to risk. Advances in free-breathing and motion-corrected T2 and T2* mapping as well as LGE imaging (120–122), without the need for breath-holding, have been already been developed to address the clinical need to further accelerate CMR data acquisition. Free-breathing, ungated, 3D contrast-enhanced CMR, which is acquired at high-resolution, is well positioned for see rapid uptake in the clinical arena for myocardial tissue characterization in the post-MI setting. Other developments in real-time cine imaging for LVEF and future refinements in CMR fingerprinting techniques (123) (for robust and fast acquisition of simultaneous T1 and T2 mapping data per slice within 1 breath-hold) will further reduce scan time and will pave the way to make CMR more accessible to a wider number of post-infarction patients. These efforts could reduce the duration of the scan to 20 min or even less, which could reduce scan costs and improve the participation of more centers in MI-related CMR research.

The possibility of performing complete CMR studies without an exogenous gadolinium-based contrast agent has been proposed. However, the prognostic role of noncontrast CMR in comparison to standard contrast-enhanced techniques warrants further investigation. Some gadolinium contrast agents (lineal contrasts, not the ones used at present) can accumulate in some tissues (124), but the clinical significance is unknown.

Edema imaging needs to be optimized for comprehensive characterization of reversible and irreversible myocardial injury after reperfusion. One of the important challenges in the field is the development of methods to differentiate intracellular from extracellular water. The role of edema as potential therapeutic target in I/R could serve as a basis for the study of myocardial water distribution using CMR (15,33).

Imaging of remote myocardium after STEMI may provide new insights into the pathophysiology of post-MI remodeling and provide incremental risk stratification value. Given the superior diagnostic performance of quantitative mapping techniques, potential novel imaging biomarkers have emerged. Specifically, recent evidence shows an abnormal tissue composition in the remote myocardium of STEMI patients (118,119,125).

In summary, CMR holds great promise to become accessible to more patients and to be used as a tool to risk-stratify patients, guide treatment, evaluate cardioprotective therapies, and improve clinical outcomes in STEMI patients. Addressing the previously mentioned questions and directions will further extend the already established role of CMR in the assessment of myocardial infarction.

**ADDRESS FOR CORRESPONDENCE:** Dr. Borja Ibanez, Centro Nacional de Investigaciones Cardiovasculares Carlos III (CNIC), c/ Melchor Fernandez Almagro, 3, 28029 Madrid, Spain. E-mail: bibanez@cnic.es. Twitter: @Borjaibanez1. OR Dr. Valentin Fuster, Mount Sinai School of Medicine, Cardiovascular Institute, One Gustave Levy Place, Box 1030, New York, New York 10029-6500. E-mail: valentin.fuster@mountsinai.org. Twitter: @CNIC_CARDIO.
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APPENDIX For supplemental material, please see the online version of this paper.