Bifunctional staining for \textit{ex vivo} determination of area at risk in rabbits with reperfused myocardial infarction

Yuanbo Feng, Zhan-Long Ma, Feng Chen, Jie Yu, Marlein Miranda Cona, Yi Xie, Yue Li, Yicheng Ni

Yuanbo Feng, Feng Chen, Jie Yu, Marlein Miranda Cona, Yicheng Ni, Theragnostic Laboratory, Department of Imaging and Pathology, Biomedical Sciences Group, 3000 KU Leuven, Belgium
Zhan-Long Ma, Departments of Radiology, the First Affiliated Hospital, Nanjing Medical University, Nanjing 210029, Jiangsu Province, China
Yi Xie, Departments of Electronics and Information System (ELIS), Ghent University, Sint-Pieersnieuwstraat 25, 9000 Gent, Belgium
Yue Li, Lab of Translational Medicine, Jiangsu Academy of Traditional Chinese Medicine, nanjing 210028, Jiangsu Province, China
Yicheng Ni, Radiology Section, University Hospitals, 3000 Leuven, Belgium

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Correspondence to: Yicheng Ni, MD, PhD, Professor, Radiology Section, University Hospitals, KU Leuven Herestraat 49, 3000 Leuven, Belgium. yicheng.ni@med.kuleuven.be

Telephone: +32-16-330165 Fax: +32-16-343765

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Abstract

AIM: To develop a method for studying myocardial area at risk (AAR) in ischemic heart disease in correlation with cardiac magnetic resonance imaging (cMRI).

METHODS: Nine rabbits were anesthetized, intubated and subjected to occlusion and reperfusion of the left circumflex coronary artery (LCx) to induce myocardial infarction (MI). ECG-triggered cMRI with delayed enhancement was performed at 3.0 T. After euthanasia, the heart was excised with the LCx re-ligated. Bifunctional staining was performed by perfusing the aorta with a homemade red-iodized-oil (RIO) dye. The heart was then agar-embedded for \textit{ex vivo} magnetic resonance imaging and sliced into 3 mm-sections. The AAR was defined by RIO-staining and digital radiography (DR). The perfusion density rate (PDR) was derived from DR for the AAR and normal myocardium. The MI was measured by \textit{in vivo} delayed enhancement (iDE) and \textit{ex vivo} delayed enhancement (eDE) cMRI. The AAR and MI were compared to validate the bifunctional staining for cardiac imaging research. Linear regression with Bland-Altman agreement, one way-ANOVA with Bonferroni’s multiple comparison, and paired t tests were applied for statistics.

RESULTS: All rabbits tolerated well the surgical procedure and subsequent cMRI sessions. The open-chest occlusion and close-chest reperfusion of the LCx, double suture method and bifunctional staining were successfully applied in all animals. The percentage MI volumes globally ($n = 6$) and by slice ($n = 25$) were 36.59$\pm$ 13.68$\%$ and 32.88$\%$ $\pm$ 12.38$\%$ on iDE, and 35.41$\%$ $\pm$ 12.25$\%$ and 32.40$\%$ $\pm$ 12.34$\%$ on eDE. There were no significant differences for MI determination with excellent linear regression correspondence ($r_{global} = 0.89; r_{slice} = 0.9$) between iDE and eDE. The percentage AAR volumes globally ($n = 6$) and by slice ($n = 25$) were 44.82$\%$ $\pm$ 15.18$\%$ and 40.04$\%$ $\pm$ 13.64$\%$ with RIO-staining, and 44.74$\%$ $\pm$ 15.98$\%$ and 40.48$\%$ $\pm$ 13.26$\%$ by DR showing high correlation in linear regression analysis ($r_{global} = 0.99; r_{slice} = 1.0$). The mean differences of the two AAR measurements on Bland-Altman were almost zero, indicating RIO-staining and DR were essentially equivalent or inter-replaceable. The AAR was significantly larger than MI both globally and slice-by-slice ($P < 0.01$). After correction with the background and the blank heart without bifunctional staining ($n = 3$), the PDR for the AAR and normal myocardium was 32$\%$ $\pm$ 15$\%$ and 35.5$\%$ $\pm$ 35$\%$, respectively.
which is significantly different ($P < 0.001$), suggesting that blood perfusion to the AAR probably by collateral circulation was only less than 10% of that in the normal myocardium.

CONCLUSION: The myocardial area at risk in ischemic heart disease could be accurately determined postmortem by this novel bifunctional staining, which may substantially contribute to translational cardiac imaging research.

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Key words: Reperfused; Acute myocardial infarction; Rabbit model; Cardiac magnetic resonance imaging; Oil-red-o dye; Iodized oil

Core tip: For developing therapeutic procedures/drugs aimed at modulating infarct size after coronary artery disease, it is important not only to measure myocardial infarction but also to know the extent of area at risk (AAR). However, determination of the AAR both in vivo and ex vivo can be challenging, and controversial. In this experiment we sought to develop a reliable method to accurately localize the culprit coronary occlusion for postmortem verification after in vivo cardiac magnetic resonance imaging and to establish a new bifunctional staining as a standard reference for ex vivo area at risk identification, which may substantially contribute to translational cardiac imaging research.

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INTRODUCTION

Acute occlusion of a coronary artery initiates an expanding process of myocardial infarction (MI) and this process ends in complete necrosis of the myocardial tissue within the perfusion bed if without a prompt therapeutic intervention. To develop therapeutic procedures and/or drugs aimed at modulating infarct size, it is very important not only to measure the MI size but also to know the extent of perfusion defect zone after coronary occlusion, which is also known as the area at risk (AAR) of myocardium[3]. Since the AAR includes both infarcted/nonviable and viable/salvageable myocardium, detection of the AAR may provide an accurate index in clinical practice and preclinical research for diagnostic, therapeutic and prognostic assessment of ischemic heart disease[6,8]. To define the AAR by both in vivo and ex vivo imaging can be challenging. Currently, the most widely used methods for in vivo AAR identification are contrast enhanced computed tomography[9] or cardiac magnetic resonance imaging (cMRI)[3], radionuclide perfusion imaging[9], and contrasted echocardiography[3]. However, determination of the AAR proves difficult and controversial in clinical cardiology[3]. The T2-weighted cMRI may highlight myocardial edema and therefore often overestimate infarct size, and sestamibi single photon emission computed tomography is limited by its low spatial resolution and radioactive nature. Compared to the AAR, MI can be evaluated in vivo by delayed enhancement (DE) cMRI, an established technique that is widely accepted in clinic to reveal acute and chronic MI in patients[7]. This technique has been validated in animal models with reperfused MI using extracellular contrast agents[8,9,11].

Experimentally, to validate in vivo imaging observations, a variety of ex vivo methods have been developed and attempted to directly visualize and quantify the AAR and MI in animal experiments, because exact postmortem detection of myocardial AAR and MI is deemed necessary as the gold reference. A few decades ago, some investigators tried to assess the AAR through postmortem intracoronary injections of colored industrial dyes[12] or barium gels[13], which stained the normal myocardium but left the AAR unstained. However, highly viscous injectants and undetermined pressure could lead to inaccurate AAR delineation. Nowadays, widespread techniques used for determination of coronary perfusion or AAR are blue dyes (typically Evans blue) and radiolabeled or colored microspheres, with which the heart is perfused at the end of experiments after re-occlusion of the coronary artery[14,15]. Triphenyltetrazolium chloride (TTC) histochimical staining is usually combined with a blue dye for simultaneous visualization of normal myocardium, AAR and MI[15]. These methods have their limitations. For instance, the most commonly used water-soluble blue dye Evans-blue may precipitate out of solution and affect the subsequent TTC staining.[16] These dyes are also prone to smearing during slicing, resulting in an ambiguous border definition[17]. The radiolabeled microspheres need the complex procedures and operators have to be exposed to radiation, in addition to the difficulty in some institutions due to unavailable nuclear facilities. The colored microspheres require tissue lyses for counting without morphometry. Therefore, such analyses still remain technically challenging.

To precisely recognize the location of previously occluded coronary branch is crucial for subsequent ex vivo calculation of AAR size. Few study detailed the technical information for accurate coronary re-ligation[18]. Given the suboptimal features of above-mentioned dyes and microspheres for visualizing the AAR in preclinical study, we sought to: (1) develop a reliable method to accurately localize the previous coronary occlusion site for postmortem re-ligation; (2) establish a new bifunctional staining as standard reference for ex vivo AAR evaluation; (3) validate the method by quantitative measurement and morphometry; and (4) compare the AAR defined by the new staining to the cMRI findings at a 3.0
T clinic magnet with histological verifications. The above goals could have been realized only when has a prior-established methodology, i.e., an occlusion/reperfusion based rabbit model of acute MI and a technical platform for pre- and postmortem imaging co-localization now become available\[19,20\]. The rabbit as a mediate size experimental animal can be easily handled, with favorable cost-effectiveness and compatibility to clinically available MRI facilities with the entire heart section includable in a standard histological glass slide. The resulted outcomes are deemed pertinent to human patients\[19,21\].

MATERIALS AND METHODS

This experimental study was carried out according to a protocol (Figure 1) that was approved by the Animal Care and Use Committee of KU Leuven, Belgium.

Preparation of red iodized oil dye

The custom-made bifunctional dye of red iodized oil (RIO) was formulated by diluting 20 mg oil-red-O dye (Sigma-Aldrich) in 100 mL of any brand of commercially available iodized oil such as Lipiodol ultra fluid (Guerbet, France), which is also known as ethiodized oil consisting of about 30%-40% iodine and poppyseed oil originally used as a radio-opaque contrast agent to outline structures in radiological investigations or in combination for transarterial chemoembolization of malignant tumors\[22,23\].

Animal model of acute reperfused myocardial infarction

A total of 9 male New Zealand white rabbits (Animal House, KU Leuven, Belgium), weighing 3.2-4.0 kg, were included in this study. Animals were housed under standard laboratory conditions. The model of acute reperfused MI was prepared by slight modification on a previously introduced method\[19\]. After sedation, endotracheal intubation and mechanical ventilation, the rabbit received intravenous (IV) injection of pentobarbital (Nembutal, Sanofi Sante Animale, Brussels, Belgium) at 40 mg/kg per hour to maintain anesthesia during the open-chest operation to dissect the pericardium and expose the left circumflex coronary branch (LCx). For making a removable coronary ligation in vivo and precise re-ligation ex vivo, we used a 1/2 circle shape triangular needle (Sutura, Inc. Fountain Valley, CA, United States) with spring eyes at the end (Figure 2A).

Two silk sutures could be easily placed through the separate eyes (Figure 2B): the thicker one (2-0) was used for the LCx ligation that could be removed for reperfusion and the thinner one (5-0) was spared for later ex vivo LCx re-occlusion in order to perform postmortem bifunctional staining (Figure 2C). The acute reperfused MI was induced by tying the thicker suture with a detachable single knot for 90 min, and LCx reperfusion was achieved by pulling the exteriorized end of suture in the closed-chest condition as detailed previously\[19\]. The ECG (Accusync® 71, Milford, Connecticut, Unites States) was monitored to confirm the formation of MI and to detect the occurrence of arrhythmia, which was treated by IV dose of Xylocaine (AstraZeneca SA, Brussels, Belgium) at 2-4 mg/kg if needed. Buprenorfine hydrochloride (Temgesic; Schering-Plough, Brussels, Belgium) of 0.216 mg was intramuscularly injected to relieve pain. All rabbits were allowed to recover and kept alive until cMRI scanning 24 h after LCx reperfusion.

Cardiac magnetic resonance imaging

The rabbit was gas-anesthetized through a mask with 2% isoflurane in the mixture of 20% oxygen and 80% room air and placed supinely in a plastic holder. Using a commercial 8-channel phased array knee coil, the rabbit was subjected to cMRI at a 3.0 T clinical MRI scanner (Trio, Siemens, Erlangen, Germany) with a maximum gradient capability of 45 mT/m. The ECG and respiration were triggered and gated by a Small Animal Monitoring System (SA Instruments, Inc. Stony Brook, New York, United States). Two ECG surface electrodes were attached to the shaved thorax skin with apparently apical pulse and the left leg. The respiration control sensor was attached on the middle of the rabbit abdomen. All data were acquired during free breathing of the animal. After determining the cardiac axes with localizers, the DE cMRI using T1-weighted 3-dimensional segmented k-space inversion recovery turbo fast low angle shot sequence was performed 20 min after intravenous administration of 0.2 mmol/kg Gd-DOTA (Dotarem, Guerbet, France). Short-axis DE cMRI was acquired to depict the presence, location, and extent of the MI. The imaging parameters were: TR: 396 ms; TE: 1.54 ms; TI: 360 ms; FOV: 240 × 180 mm²; FA: 15°; and in-plane resolution: 1.1 × 0.8 mm². The ex vivo higher resolution DE-cMRI was also obtained immediately after animal sacrifice using the same MRI unit with a wrist coil as previously described\[19\]. The DE cMRI using an extracellular nonspecific contrast agent has been ap-
plied in both animal MI models and clinical patients as a gold standard technique for identification and quantification of the MI\textsuperscript{[10,11]}. Therefore, the final MI size was also defined by in vivo and ex vivo DE-cMRI in this study.

**Postmortem bifunctional staining**

Bifunctional staining of the RIO dye refers to: (1) visualization, location and delineation of the AAR by using its red staining function for further morphometric quantification; and (2) microangiographic examination of the AAR by using the digital radiography (DR) function to compare the perfusion density rate (PDR) between the AAR and normal myocardium. This staining served as a postmortem index for regional myocardial blood flow. Briefly, after the last in vivo cMRI, the rabbit was IV injected with Heparin at 300 IU/kg to prevent coagulation. Under deep anesthesia and thoracotomy, the heart was excised with a section of the aortic arch isolated and pericardial tissue trimmed. As shown in Figure 3A, a plastic catheter filled with 0.9% saline was inserted into the aorta and anchored with its tip 1 cm above the aortic valves. Care was taken to avoid air bubbles entering the catheter that was connected to a 10 mL syringe fixed on the injection bump (BD PILOT, Becton Dickinson Infusion System, France). Under a perfusion pressure of 100 mmHg, 10 mL saline was infused until the eluate from the coronary sinus became clear. The heart was rinsed to wash out remaining blood. Then, the 5-0 suture spared at the coronary occlusion site during open-chest operation was tightened for causing re-occlusion, and 4 mL of RIO dye (Figure 3B) was infused through the aortic catheter for 15 min under the same pressure by using the same pump as mentioned above. The infusion was stopped when the normal myocardium was completely stained red by the RIO dye (Figure 3C). Upon this perfusion staining technique with the LCx re-ligated, the normal myocardium stained brick red and the AAR remained uncolored. Afterwards, the heart was imbedded in agar solution for ex vivo cMRI, followed by serial sectioning into 6-8 slices of 3.0 mm thickness to match with the corresponding short axis cMRI images. Such stained heart sections were exposed by DR with a mammographic unit (Embrace; Agfa-Ge-vaert, Mortsel, Belgium) at 25 KV and 18 mA, and photographed by a digital camera before and after 24 h of fixation with 10% formalin. The fixed heart sections were imbedded into paraffin and processed for histology with 5 micron thick hematoxylin-eosin (HE) stained slices for microscopic inspection.

**Imaging analysis**

The MI size on cMRI was determined in consensus by three authors with the built-in software of Siemens workstation (SyngoMR A30). To facilitate cardiac morphometric comparisons with different imaging methods, the dimension was calculated as a value relative to the entire left ventricular myocardium. The MI size (%) on DE cMRI was derived semi-automatically by computer to count all enhanced pixels with the signal intensity higher than two standard deviations above the mean of image intensities in the remote NM, which were divided by all ventricular pixels on the same short-axis image. Digital photographs of RIO-staining sections and DR images
were transferred to a personal computer for planimetric analysis by using Image J 1.38x software (Research Services Branch, NIH, Bethesda, MD, United States). The AAR defined as non-stained area by RIO-dye and non-opaque area by DR was contoured with consensus on the computer screen. Due to partial volume effect and fatty tissue interference on DE cMRI, the apical and basal slices were excluded and the remaining slices ($n = 25$) obtained from 6 rabbits were used for the comparisons of the corresponding bifunctional staining sections.

Perfusion density rate measurement
The measurement of perfusion density rate (PDR%) measurement was performed on DR images of the heart sections using the same Image J 1.38x software. The blank myocardial density refers to the DR density of the heart tissues from control group ($n = 3$) without RIO dye staining. The AAR of low density and the normal myocardium of high density were delineated in all sections ($n = 25$) with an operator-defined region of interest. The mean gray value (GV) of all sections were automatically generated and normalized by the background GV. PDR% refers to the increased perfusion density relative to background, which was calculated by the following formula: $\text{PDR\%} = \frac{(\text{GV}_{\text{region of interest}} - \text{GV}_{\text{blank myocardial density}})}{\text{GV}_{\text{background}}} \times 100$.

Statistical analysis
Numerical data were expressed as mean ± SD. Comparisons among multiple factors were performed using one way ANOVA, followed by the Bonferroni’s test to identify any differences between each two techniques. All tests were two tailed. The correlations were tested by Spearman’s method. The linear regression and the means of Bland-Altman analysis were used for correlation and agreement, respectively. Correlations were described by the correlation coefficient with 95% CIs. A difference was considered statistically significant if the P value was less than 0.05. The paired student $t$ test was used for comparison of the PDR between the AAR and MI. The statistical analyses were performed with Analyse-it® 2.14 for Excel and GraphPad Prisma® 5.0.

RESULTS
General outcomes
All animals survived the anesthesia, surgical procedures for inducing acute reperfused MI and subsequent cMRI. As one of the critical steps, the double suture method was successfully applied in all animals, which ensured later bifunctional staining. The normal myocardium from all rabbits was well perfused by the RIO dye. Figure 4 demonstrated a case with a large reperfused MI by different techniques. The hyperenhanced transmural MI appeared similarly on both iDE and eDE (Figure 4A, B), which was somewhat smaller than the AAR as an unstained region by RIO (Figure 4C) and non-opacified region on DR (Figure 4D). With this bifunctional staining, the large haemorrhagic MI appeared brown greyish (Figure 4C) after overnight Formalin fixation. The myocardial necrosis
Figure 4  An example of a reperfused hemorrhagic myocardial infarction (arrow) shown by cardiac magnetic resonance imaging and the area at risk demonstrated with bifunctional staining method. A: The hyperenhanced transmural myocardial infarction (MI) was seen at the lateral wall on in vivo delay enhancement (DE) cardiac magnetic resonance imaging (cMRI); B: The ex vivo DE cMRI was highly correspondent with A; C: In contrast to the normal myocardium with red staining, the area at risk (AAR) was shown as an unstained region, on which the brown grayish area indicates a large intramural hemorrhagic infarction after the fixation by Formalin; D: On digital radiography (DR), the AAR was shown as non-opacified region in contrast to the opacified normal myocardium. Notice that the AAR defined by red iodized oil-staining (RIO-staining) in C and DR in D was somewhat larger than the MI defined by in vivo DE cMRI in A and ex vivo DE cMRI in B; E: The hematoxylin-eosin (HE) stained macroscopic view was obtained from the corresponding histological section of A, B, C, and D; F: Photomicroscopic view of the HE stained slice (magnification, ×100) confirmed the presence of myocardial necrosis with hemorrhage and tissue reaction. h: Hemorrhagic infarction; n: Adjacent normal myocardium with inflammatory infiltration.

with haemorrhage was finally confirmed by macro- and microscopic views of HE stained slice (Figure 4E, F).

**Magnetic resonance imaging findings**
The location and extent of the MI were clearly shown as the hyperenhanced region on DE cMRI in all cases. The percentage MI sizes globally ($n = 6$) and slice-by-slice ($n = 25$) were 36.59% and 35.41% on in vivo DEcMRI (iDE), and 32.88% and 32.40% on ex vivo DEcMRI (eDE). There was no significant difference between iDE and eDE. Linear regression analyses showed an excellent correlation both globally and slice-by-slice between MI volumes on iDE and eDE ($r = 0.89, 0.98$) (Figure 5A, B). By Bland-Altman analysis, there was a 1.2% bias with 95%CI of 2.4% to 4.7% in global heart (Figure 5C) and 0.5% bias with 95%CI of 2.6% to 3.6% by slice (Figure 5D), indicating high agreement of the two measurement methods.

**Bifunctional staining findings with red-iodized-oil dye**
The RIO dye was successfully formulated (Figure 3B). It showed homogenous distribution into the accessible cardiac tissues. After staining, the normal myocardium appeared scarlet red (Figure 3C) and rose red (Figure 4C) before and after formalin fixation and as more opacified region on DR images (Figure 4D), whereas the AAR showed as non-RIO-stained region on heart slices (Figure 4C) and the less opacified region on DR image (Figure 4D). By bifunctional staining, the percentage AAR$_{global}$ was 44.8% ± 15.18% and 44.74% ± 15.98%, and percentage AAR$_{slice}$ was 40.04% ± 13.64% and 40.48% ± 13.26% by RIO-staining and DR, respectively. There was no significant difference between RIO-staining and DR ($P = 0.9$). Linear regression analyses showed an excellent correlation both globally and slice-by-slice between AAR volumes defined by RIO-staining and DR ($r = 0.99, 1.0$) (Figure 6A, B). The limits of Bland-Altman agreement were 3.2%-3.0% (Figure 6C) by slice and 2.3%-1.8% (Figure 6D) in global measurements with RIO-staining and DR respectively. The mean difference was almost zero, indicating a very high agreement between RIO-staining and DR techniques.

**Comparing different techniques**
The MI quantified from both iDE and eDE cMRI were significantly smaller than the AAR$_{RIO-staining}$ and AAR$_{DR}$ with bifunctional staining ($P < 0.01$; Figure 7), confirming that the final MI was smaller than ischemic region.
However, the MI on eDE-cMRI correlated well with AAR<sub>RIO-staining</sub> and AAR<sub>DR</sub> (Figure 8), suggesting that the bifunctional staining with RIO dye could successfully delineate and characterize the AAR.

**Perfusion density rate**

After aortic infusion with the RIO dye while the LCx was re-ligated, the PDR of the normal myocardium and AAR became 354.89% ± 35.18% and 32.28% ± 15.06% relative to the equipment background. The PDR was significantly lower in the AAR, i.e., only about 9.10% ± 3.93% of the PDR in the normal myocardium ($P < 0.001$) (Figure 9), suggesting insufficient collateral coronary blood supply during the 90 min occlusion, leading to the rabbit model of reperfused MI.

**DISCUSSION**

We studied myocardial AAR in the rabbit heart after 90 min of LCx occlusion followed by reperfusion. The AAR refers to the myocardium without blood supply during coronary artery occlusion. The final MI was detected by *in vivo* and *ex vivo* DE cMRI. The main outcomes of the present study are: (1) the newly developed method of bifunctional staining enabled reliable high-quality images; (2) both *in vivo* and *ex vivo* DE cMRI allowed visualization and quantification of the MI; (3) though measured with different methods, the dimension of the AAR was significantly larger than that of the MI, suggesting the potential of this method to help discriminate the well perfused normal myocardium from the non-perfused nonviable and/or less perfused by viable probably salvageable myocardium; and (4) the PDR was 10 times lower in the AAR compared to the normal myocardium, suggesting that the bifunctional RIO dye may facilitate not only morphometry but also perfusion quantification.

The animal models of reperfused MI have been extensively applied for cardiovascular research both in large and small animals. How to accurately re-ligate coronary artery branch to replicate the exact occlusion site prior to reperfusion appears crucial. The majority of the authors used the snare-loop method for coronary artery occlusion-reperfusion on the open-chest MI model with later postmortem vessel re-ligation<sup>[24]</sup>. Other studies described re-oclusion of coronary branches with a balloon catheter under the close-chest conditions<sup>[25]</sup>. However, there is a lack of detailed information about how to imitate postmortem the exact *in vivo* artery occlusion, which
is though a prerequisite for defining the AAR or ischemic area. We have previously introduced an effective method for \textit{in vivo} LCx occlusion and reperfusion\textsuperscript{[19]}. In this study we modified that method by using the double sutures to precisely co-localize LCx obstruction both \textit{in vivo} and \textit{ex vivo}. Within the same puncture, one suture was for \textit{in vivo} coronary occlusion and reperfusion, and the other suture was left spared \textit{in vivo} but tightened with a knot to re-

Figure 7  Comparisons between myocardial infarction and area at risk by different techniques. There were no significant differences between myocardial infarction \textit{in vivo} delayed enhancement (MI iDE) and myocardial infarction \textit{ex vivo} delayed enhancement (MI eDE) measured from cardiac magnetic resonance imaging in global (left) and slices (right). There were no significant differences between area at risk digital radiography (AAR DR) and area at risk red iodized oil-staining (AAR RIO-staining) in global (left) and slices (right). The significant differences were observed on AAR DR vs MI iDE ($P < 0.01$); AAR DR vs MI eDE ($P < 0.01$); AAR RIO-staining vs MI iDE ($P < 0.01$); AAR RIO-staining vs MI eDE ($P < 0.01$). NS: Not significant.

Figure 6  The linear correlations (above) and Bland-Altman analyses (bottom) to compare the area at risk by red iodized oil-staining and digital radiography of the bifunctional staining. The plots showed a good correlation between the area at risk (AAR) measured with red iodized oil (RIO)-staining and digital radiography (DR) on global (A) and slices (B). Plots of the differences in AAR on global (C) and slices (D) were almost zero, indicating that RIO-staining and DR are essentially equivalent.

![Scatter plot with fit](image1)

![Scatter plot with fit](image2)

![Difference plot](image3)

![Difference plot](image4)
occlude the coronary branch \textit{ex vivo}. This double suture method was successfully validated in all animals, which is a critical step to ensure later bifunctional staining to obtain the desired results as presented.

In the cardiac imaging research on the ischemic heart disease using animal hearts, a most crucial issue is adequate staining of the AAR and discrimination between risk and non-risk zones, because a correctly defined borderline between the ischemic and necrotic zones will have a direct impact on MI diagnosis and therapeutic interventions. In this article we described a new bifunctional staining method to define the AAR, which, to our best knowledge, has not been reported in the literature.

Function 1 is to visualize the normal myocardium in red color but the AAR uncolored utilizing the oil-red-o component of the custom-made RIO dye, which facilitates morphometric delineation, quantification and comparison; whereas function 2 is to enable visualization and volumetric density measurement of the non-opacified normal myocardium and the opacified AAR by utilizing the iodized oil component of the RIO dye for making DR to extrapolate, quantify and evaluate the difference in hemodynamic features such as coronary vessel density, interstitial distribution space and/or regional myocardial blood flow/volume between the normal and ischemic myocardium.

There have been some methods reported in the literature for postmortem delineation of the AAR. Coronary infusion of hydrophilic Evans-blue solution is the most effective method to differentiate the AAR region from the normal myocardium. According to our best knowledge, the RIO dye utilized in our study is the first bifunctional red-iodized oil staining method that allows differentiation of the AAR from normal myocardium on digital radiography.

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\begin{figure}[ht]
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\includegraphics[width=\textwidth]{fig8}
\caption{The linear correlations between \textit{ex vivo} cardiac magnetic resonance imaging and bifunctional staining. The myocardial infarction (MI) measured by global (left) and slices (right) on \textit{ex vivo} delayed enhancement cardiac magnetic resonance imaging corresponded well with area at risk by red iodized oil-staining (above) and digital radiography (bottom), confirming the bifunctional staining with RIO dye could successfully delineate and characterize the AAR. MI<sub>eDE</sub>: Myocardial infarction \textit{ex vivo} delayed enhancement; AAR<sub>RIOSTAINING</sub>: Area at risk on red-iodized oil-staining; AAR<sub>DR</sub>: Area at risk on digital radiography.
}
\end{figure}

\begin{figure}[ht]
\centering
\includegraphics[width=\textwidth]{fig9}
\caption{Comparison of perfusion density rate between area at risk and normal myocardium. Coronary artery occlusion caused a significant decrease of blood perfusion in ischemic region compared to non-ischemic region ($P < 0.001$), resulted in a difference over 10 times in-between. AAR: Area at risk.
}
\end{figure}
common technique to stain the remote myocardium while the insulted coronary branch is re-ligated\[23-27\]. Evans-blue is a water-soluble intravital dye with strong albumin binding capacity and has been used in hemodynamic studies for decades\[28-29\]. More recently Evans-blue has been found with high necrosis avidity and used to clearly mark the well-reperfused necrotic tissues\[30-32\]. Therefore, there is a risk of underestimation of the AAR due to contamination of the inwards diffused Evans-blue to the ischemic necrosis\[33\]. Some other groups tried to use TTC for demarcating the AAR, however TTC perfusion induced severe tissue contraction, which may affect the geometry of heart and might prevent subsequent successful dye perfusion\[34\]. Other studies used fluorescein dyes under dark background or radio- and color-labeled microspheres to define or quantify the AAR\[33-35\]. However, all these methods are only unifunctional and/or lacking required accuracy. The custom-made bifunctional RIO dye as introduced in this study appears a much better alternative to Evans blue. The first component of RIO dye is oil-red-O, which is a stable lipophilic pigment and can be easily diluted in any lipid media such as the iodized oil as the second component of this dye. The distribution of the RIO dye in the cardiac tissue after intracoronary infusion seems only pressure dependent rather than being diffusible afterwards driven by its concentration gradient due to the fact that a hydrophobic dye exists in the hydrophilic environment of myocardium, resulting in a more accurate delineation of the borders of the AAR without infiltration of the dye into the adjacent ischemic region.

Suspension of barium sulphate has also been used for postmortem coronary angiography\[13,38\]. However, the heavy particles tended to precipitate in its water media, leading to suboptimal perfusion to vascular bed with smaller vessels. As shown in this study, the second component of iodized oil in the RIO dye solved the problem well in DR and its quantification. Thus, the golden combination of the two dye components facilitated successfully the bifunctional staining and enabled comprehensive AAR study.

We have gained the following experiences: (1) hepatinizing the animal prior to sacrifice and rinsing the coronary bed with normal saline beforehand proved beneficial for the next bifunctional staining of the AAR by RIO dye perfusion; (2) controlling adequate pressure and observing epicardial red coloration during dye infusion were crucial for the success of procedure; and (3) fixing the heart sections with formalin overnight helped color differentiation between the normal myocardium (rose red) and AAR (gray or brown grayish depending on the degree of hemorrhage) and later planimetric quantification due to a clear borderline.

Myocardial blood volume (capillary bed) and myocardial blood flow can be determined by microsphere methods in animals after excision of the heart\[39,40\]. Such methods, however, require cutting the myocardium into discrete samples, which may be a combination of normal tissue, border zone, hyperemic tissue, and severely ischemic tissue. Thus, the processing of selected tissue samples can bias the results as well as blur the exact relation to overall myocardial morphology. Although the degree of resolution for flow measurement could possibly be improved by cutting myocardium into smaller samples, this becomes tedious and impractical for studying flow throughout the heart. Besides, it is more difficult to get topographic information.

Using the PDR derived from microangiography on DR to indirectly evaluate vascular blood perfusion to viable and necrotic tissues has been reported\[41\]. In our experiment, The PDR was significantly lower in AAR, which was about 9% of the normal myocardium. This finding is close to the results of Connelly’s group on myocardium blood flow measured by radioactive microspheres on the similar animal setting\[42\]. The other group showed that in the reperfusion MI dog model, myocardial samples with only mild (about 20%-30%) reductions in myocardial blood flow compared to non-ischemic regions\[43\], which appears higher than our results probably due to the richer collaterals of coronary artery in dogs than in rabbit\[40\]. The application of this new staining method on our rabbit model was successful and can be translated to other species.

Accurate quantitative measurement of MI size is an important aspect of cardiovascular research. The in vivo cMRI for accurate evaluation of MI has been described previously in rabbit models\[44\]. In this study we have performed the high resolution ex vivo cMRI with extensive postmortem evaluations on rabbit hearts, which were supported by histological staining and consistent to in vivo cMRI. The bifunctional staining could be incorporated into the study using clinically available MRI facilities to acquire outcomes that are more pertinent to human patients.

**Study limitations**

First, the AAR was demarcated by intracoronarily infusing the heart with the RIO dye after local re-occlusion. Such a procedure was time-consuming with possible technical failure to obtain homogeneous staining of all accessible normal myocardium. Secondly, the postmortem dye distribution may differ from in vivo blood perfusion of the normal myocardium when the related artery is occluded, because normal coronary flow is influenced by the intraventricular pressure so the flow in postmortem infusion may not exactly simulate the in vivo flow pattern. Furthermore, in this experiment by using RIO dye, we could achieve accurate delineation of the AAR postmortem, but we could hardly discriminate the MI from any ischemic but still viable tissues within the AAR. Considering these drawbacks, we have to further optimize this experimental setting for determination of both AAR and MI under both in vivo and ex vivo conditions.

In conclusion, this article introduced a novel bifunctional staining method to accurately visualize and quantify AAR postmortem, which was successfully applied in rab-
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