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MINI-FOCUS: INFLAMMATION IN CARDIAC INJURY

Acute CD47 Blockade During Ischemic Myocardial Reperfusion Enhances Phagocytosis-Associated Cardiac Repair

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HIGHLIGHTS

- New therapies are needed to enhance myocardial salvage after myocardial ischemia and reperfusion.
- Human and murine hearts express CD47 and calcretinin that increased after ischemia and reperfusion.
- Phagocytic efficiency of dying cardiac myocytes was enhanced after antibody-mediated blockade of either myocyte CD47 or macrophage CD47-ligand, SIRPα.
- After ischemia and reperfusion, enhancement of dead myocyte clearance by macrophages, after CD47 blockade, improved inflammation resolution, reduced infarct size, and preserved cardiac systolic function.

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SUMMARY

Our data suggest that, after a myocardial infarction, integrin-associated protein CD47 on cardiac myocytes is elevated. In culture, increased CD47 on the surface of dying cardiomyocytes impairs phagocytic removal by immune cell macrophages. After myocardial ischemia and reperfusion, acute CD47 inhibition with blocking antibodies enhanced dead myocyte clearance by cardiac phagocytes and also improved the resolution of cardiac inflammation, reduced infarct size, and preserved cardiac contractile function. Early targeting of CD47 in the myocardium after reperfusion may be a new strategy to enhance wound repair in the ischemic heart. (J Am Coll Cardiol Basic Trans Science 2017;2:386–97) © 2017 The Authors. Published by Elsevier on behalf of the American College of Cardiology Foundation. This is an open access article under the CC BY-NC-ND license (http://creativecommons.org/licenses/by-nc-nd/4.0/).

Although clinical management of acute myocardial infarction (AMI) has significantly reduced morbidity and mortality, the consequence of these advances include an emerging incidence of post-MI heart failure (1). In turn, new approaches that enhance cardiac wound healing and are complementary to the current standards of care have the potential to improve on left ventricular (LV) systolic function (2).

A critical determinant of heart failure susceptibility after AMI is infarct size (3). Infarct size and loss of nonregenerative cardiomyocytes (CMs) by acute necrosis directly correlates with ventricular dysfunction and heart failure (4). Infarct necrosis can also expand during reperfusion injury or after maladaptive inflammation, leading to accelerated and adverse ventricular remodeling (5,6).

During wound healing, clearance of dying cells must occur efficiently to prevent secondary necrosis and prolonged inflammation. Efficient phagocytic clearance of apoptotic cells by macrophages (Mφs) via efferocytosis, actively programs cellular inflammation-resolution and tissue repair signaling pathways (7). This clearing is particularly important in the heart, where inefficient removal of necrotic CMs may lead to collateral myocyte loss and infarct expansion (5,8). In a previous study, we experimentally linked the efficiency of phagocytic clearance to infarct size expansion and discovered that Mφs defective for efferocytosis led to deteriorated cardiac function after MI (9). These studies begged additional questions, including the natural efficiency of CM clearance by phagocytic cells and whether strategies that target phagocytic enhancement might enhance heart healing. In this study, we focused on CM-intrinsic factors, namely CD47, that regulate phagocytic interactions with Mφs (10).

Dying cell engulfment requires cell surface presentation of so-called “eat-me” signals, such as phosphatidylserine and calreticulin, which must occur in tandem with downregulation of anti-phagocytic “don’t-eat-me” markers, including CD47 (10). Independent of programmed cell death per se, blockade of CD47 and concomitant overexpression of calreticulin can permit phagocyte ingestion of viable cells. In addition to experimental blockade (11), natural reduced expression of CD47 occurs in vivo, for example, in senescent erythrocytes; this is associated with red blood cell erythrophagocytosis and clearance from the circulation by resident tissue Mφs (12). A recent study published in the journal Nature (13) highlights the potential of CD47 blocking strategies in atherosclerotic cardiovascular disease.

Although CD47 expression has been profiled in neonatal CMs (14), much less is known about its expression and function in adult CMs. CD47 deficiency is associated with enhanced cardiac performance after administration of vasoactive agents (15). In skeletal muscle, CD47 can regulate PGC-1α-dependent mitochondrial biogenesis through recognition of its thrombospondin ligand (16). CD47 has also been linked to signaling in smooth muscle cells (17). The significance of CD47 after AMI is unclear.

Our preliminary mechanistic studies suggested elevated CD47 expression in CMs relative to other cardiac cells, leading us to hypothesize that CMs may exhibit natural resistance to phagocytic removal after injury. To test in principle whether acute CD47 blockade during reperfusion of ischemic myocardium enhances cardiac repair, in association with CM phagocytosis, we used an experimental MI model in mice and injected anti-CD47-blocking antibodies during reperfusion. Importantly, CD47 targeting was limited to a single time and dose to focus actions during the height of CM phagocytosis and to avoid targeting alternative postacute CD47 signaling pathways.

ABBREVIATIONS AND ACRONYMS

IAP = integrin-associated protein
AMI = acute myocardial infarction
CM = cardiomyocyte
LV = left ventricular
Mφ = macrophage
NO = nitric oxide

MATERIALS AND METHODS

HUMAN SAMPLES. Human cardiac samples were obtained from the Northwestern University Department
of Pathology and the office of the Medical Examiner of Chicago. Under the protocol approved by the institutional review board at Northwestern University (#STU00079445), archived formalin-fixed paraffin-embedded cardiac autopsy samples were obtained and analyzed at Northwestern University Feinberg School of Medicine, as previously described (18). Specifically, autopsy cases were evaluated by a combination of factors, including serum cardiac enzymes and gross evidence at time of autopsy. Gross evidence consisted of pale or yellow myocardial areas with or without a hyperemic border. Age of infarct ranged from 6 h to several days. Time from death to autopsy ranged from 18 h to 7 days. Autopsy tissue blocks of myocardium from acutely infarcted areas versus noninfarcted areas and noninfarcted individuals were fixed in 10% formalin and serial 6-μm sections were stained with hematoxylin and eosin. In this study, a total of 22 human cardiac autopsy samples were used, 11 from AMI cases and 11 from non-AMI cases.

**Murine Samples.** Murine hearts were perfused and fixed with 10% phosphate-buffered formalin at physiological pressures. Hearts were cut transversely, parallel to the atrioventricular groove and coronary sulcus. Fixed hearts were embedded in paraffin at the Northwestern University Mouse Histology and Phenotyping Laboratory. Blocks were serially cut 6 μm apart. Alternating sections were stained with hematoxylin and eosin. For frozen sections for immunohistochemistry, samples were rinsed and incubated overnight in 7% sucrose and frozen in freezing medium unless otherwise indicated. Transverse cryosections were cut at a thickness of 10 μm on a Leica cryostat and placed on super frost plus-coated slides for analysis.

**Mice.** B6; D2-Tg(Myb6*+mCherry)2Mik/J mice were from Jackson Laboratory (stock No: 021577, Bar Harbor, Maine). The alphaMHC-mCherry transgene has a modified mouse alpha myosin heavy chain promoter sequence directing cardiac-specific mCherry expression in CMs. Mice were housed in temperature- and humidity-controlled environments and kept on a 12:12-h day-night cycle with access to standard mouse chow and water ad libitum. All studies were approved and reviewed by the Institutional Animal Care and Use Committee at Northwestern University (Chicago, Illinois). CD47+/− mice were obtained from the Jackson Laboratory (stock #003173).

**Materials and Antibodies.** Tissue culture dishes were from Corning (Corning, New York), and fetal bovine serum was from Gibco (Gaithersburg, Maryland). Chemical reagents were from Sigma Chemical Co. (St. Louis, Missouri), unless stated otherwise.

**Antibodies.** CD47 monoclonal and polyclonal antibodies were as previously described (19). Anti-CD68 was obtained from Abcam (ab125212, Cambridge, United Kingdom). Monoclonal mouse anti-Desmin (clone DE-U 10) from mouse ascites fluid was from Sigma (D1033). Rat antimouse Mφ antibody Mac-2 was from Cedarlane (Burlington, Ontario, Canada). Rat antineutrophil antibody was from Serotec (Raleigh, North Carolina).

**Primers for Semi-Quantitative and Real-Time PCR in This Study.**

<table>
<thead>
<tr>
<th>Primer</th>
<th>Sequence</th>
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<tr>
<td>cd2f</td>
<td>TCT GGA TCG GAA CCA AAT GA</td>
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<tr>
<td>tf</td>
<td>GAGATACACTCTCCACA-CAGACC</td>
</tr>
<tr>
<td>tnr</td>
<td>CTT ACG ATC TTC GTT CAT ACA</td>
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<tr>
<td>il-10</td>
<td>GCC AAG CCT TAT CGG AAA TG</td>
</tr>
<tr>
<td>hcd47f</td>
<td>5'-AGC TCT AAA CAA GTC CAC TGT CCC-3'</td>
</tr>
<tr>
<td>hcd47r</td>
<td>5'-TCC TGT GTG TGA GAC AGC ACT-3'</td>
</tr>
<tr>
<td>control</td>
<td>TGC CTA GAC GTG CTT GCA</td>
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<tr>
<td>htp</td>
<td>5'-AGC TCT AAA CAA GTC CAC TGT CCC-3'</td>
</tr>
<tr>
<td>htpb</td>
<td>5'-TGA GTT CAT ACC GTG CTG CTA-3'</td>
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**Ischemia and Reperfusion.** Ischemia and reperfusion injury was performed in male and female B6 mice 8 to 12 weeks of age. Ischemia was produced by occluding the left coronary artery with a 7-0 silk suture on a tapered tube for 45 min as we have previously described (20) followed by reopening of the ligature to allow for reperfusion. Mice were anesthetized with tribromoethanol (Avertin) and received buprenorphine (0.1 mg/kg subcutaneously) before surgery and after survival surgery, every 12 h up to 48 h for pain management.

**Laser Capture Microdissection.** RNA from myocardial sections was captured by laser-capture microdissection using a Zeiss P.A.L.M. laser microdissection system as previously described (21). Total RNA was isolated using the RNAqueous-Micro kit from Ambion (Waltham, Massachusetts) and reverse-transcribed into cDNA using SuperScript III First-Strand Synthesis Mix (Invitrogen, Carlsbad, California).

**Semi-Quantitative and Quantitative Reverse Transcriptase Polymerase Chain Reaction.** Hearts were snap-frozen for RNA. Total RNA was extracted using the RNeasy kit (Qiagen, Hilden, Germany). cDNA was synthesized from 4 μg of total RNA using oligo (dT) and Superscript II (Invitrogen). cDNA was subjected to quantitative reverse transcriptase polymerase chain reaction amplification using a SYBR Green PCR Master Mix (Applied Biosystems, Foster City, California).
ADMINISTRATION OF ANTI-CD47 VIA INTRAMYOCARDIAL DELIVERY. Five minutes before the ligature was released, 100 µg anti-CD47 (mIAP301) monoclonal antibody was injected at 3 sites into the ischemic myocardium, just distal to the coronary ligation with 30 µl of antibody solution per site. At the end of 45 min of occlusion, the ligature was released, and the heart was reperfused.

ADENOVIRUS ADMINISTRATION. CM specific-CD47 expression was induced as previously described (22). Adeno-associated virus1-cTNTP-GFP, 2A-mCD47-WPRE and adeno-associated virus1-cTNTP-GFP were from Vector Biolabs (Malvern, Pennsylvania). We administered 1 x 10^12 adeno-associated virus into 6-week-old B6 mice and hearts were harvested for immunohistochemical staining to check for increased protein.

FLOW CYTOMETRY PREPARATION (CARDIAC). Flow cytometric analysis of cells after MI. Mice were anesthetized with isoflurane after MI. Peripheral blood was drawn via retro-orbital bleed with citrate solution (100 mmol/l Na-citrate, 130 mmol/l glucose, pH 6.8), as an anticoagulant. Spleens were removed, triturated in Hank's Balanced Salt Solution (Mediatech, Inc., Manassas, Virginia) at 4°C with the end of a 3-mI syringe and filtered through a nylon mesh. Cell suspension was pelleted and red blood cells were lysed with ACK lysis buffer and resuspended in flow cytometry buffer.

For infarct tissue. Hearts were harvested, perfused with saline to remove peripheral cells, minced with fine scissors, and placed into a cocktail of collagenase and DNase (Sigma-Aldrich, and Worthington Biochemical Corp., Lakewood, New Jersey) and shaken at 37°C for 1 h. Cells were triturated through nylon mesh or 70 µm strainer and centrifuged at 15 min at 500 x g and 4°C. Total cell numbers were determined by Trypan blue staining. The resulting single-cell suspensions were rinsed with Hank's Balanced Salt Solution supplemented with 0.2% (wt/vol) bovine serum albumin and 1% wt/vol fetal calf serum. Flow cytometry was performed as previously described (9) and with indicated antibodies in figure legends.

CARDIAC INFARCT SIZE MEASUREMENT. At time of humane killing, mice were anesthetized with a ketamine hydrochloride and xylazine hydrochloride solution C-IIIN from Sigma (K113). The abdominal wall below the rib cage was opened and the diaphragm was cut after lifting the sternum with tweezers. The lower part of rib cage was removed to expose the heart for removal. Fluorescent latex beads were perfused retrograde to determine the area at risk. Subsequently, hearts were sliced into 1-mm transverse cross sections and then incubated with 1% 2,3,5-triphenyl-tetrazolium chloride solution (Sigma T8877) in saline for 15 min at 37°C. Hearts were then placed in 10% formalin. Viable tissue stained red and infarcted tissue white. Heart sections were weighed. Digital photomicrographs were taken and images of infaracts were blinded for analysis. Infarct size was determined as a percentage of the left ventricle. Infarct area and the total area of LV myocardium were traced manually in the digital images. Infarct size, expressed as a percentage of the area at risk, was calculated by dividing the sum of infarct areas from all sections by the sum of LV areas from all sections and multiplying by 100.

GADOLINIUM-ENHANCED MAGNETIC RESONANCE IMAGING. Digital photomicrographs were taken and images of infaracts were blinded for analysis. Infarct size was determined as a percentage of the left ventricle. Infarct area and the total area of LV myocardium were traced manually in the digital images. Infarct size, expressed as a percentage of area at risk, was calculated by dividing the sum of infarct areas from all sections by the sum of LV areas from all sections and multiplying by 100. Magnetic resonance imaging was performed on a 7-T Clinscan system (Bruker, Ettingen, Germany) equipped with actively shielded gradients (BGA12) with a full strength of 440 mT/m and a slew rate of 3,440 mT/m/ms. Images were acquired using a receive only 4-channel phased array radiofrequency coil (using a body coil for radiofrequency transmission) and an magnetic resonance-compatible physiological monitoring and gating system for mice (SA Instruments, Inc., Stony Brook, New York). The magnetic resonance imaging was performed on day 1 after the MI. The magnetic resonance imaging protocol included multislice localizer imaging to select a short axis LV slice. Late gadolinium imaging was performed using a cardiorespiratory gated multislice inversion recovery sequence covering the entire LV from base to apex. Typical imaging parameters included TE/TR, 2.3/4.7 ms; slice thickness, 1 mm; number of slices, 7 to 8; inversion time, 550 ms; number of averages, 2; spatial resolution, 0.2 x 0.2 mm^2; flip angle, 300; and scan time, 7 min. Gadolinium (0.5 mmol/kg body weight) was injected intraperitoneally and late gadolinium enhancement imaging was started 20 minutes after the injection. Image analysis was performed on a workstation using Segment software (Medviso, Lund, Sweden). Using the software, epicardial and endocardial contours were drawn and Otsu thresholding method was used to automatically
delineate the infarct zone. Infarct size was calculated as a percentage of the total LV mass.

**CARDIAC EFFEROCTYSIS ASSAYS.** Transgenic Myh6-driven mCherry mice underwent the surgery as described and flow cytometry of myocardial extract was performed after the MI to identify phagocyte, mCherry double positive cells, as previously described (18). Cells were trypsinized to dissociate cell-cell interactions and reveal only internalized mCherry signal. Cardiac phagocytosis was also measured in situ by counting the number of free and ingested apoptotic cells in individual sections. Apoptotic cells were considered “free” when they were not surrounded by, or in contact with, Mφs. The analysis was performed in a blinded fashion by 2 independent observers.

**ECHOCARDIOGRAPHY.** Two-dimensional transthoracic echocardiography was performed using a 25-MHz probe (Vevo 770, Visualsonics, Toronto, Canada) with mice in a supine position. The mouse chest was treated with a depilatory agent and mice were anesthetized with isoflurane (2% in O2) and heart rate was monitored. Left ventricle dimensions were assessed on Visualsonics software in a short axis and long axis views from the mid-left ventricle, just below the papillary muscles. In short axis, 2D (B-mode) images were taken every millimeter starting from apex to base; subsequently, an M-mode was taken 1 mm before, at, and after the papillaries. All measurements were made in 2 to 6 consecutive cardiac cycles and the averaged values used for analysis. The LV end-diastolic and end-systolic dimensions and the thickness of the interventricular septum and posterior wall were made from the M-mode tracings and fractional shortening (ratio between diameter of the LV in diastole or relaxed versus diameter when contracted) was also measured as an indicator of systolic function. Dimensions were measured between the anterior wall and posterior wall. Diastolic measurements were made at the point of minimal cavity dimension, using the leading edge method of the American Society of Echocardiography.

**CELL TYPES.** For isolation of primary Mφs, when indicated, bone marrow-derived Mφs were isolated from murine femurs and cultured in L-cell-conditioned medium, containing macrophage colony-stimulating factor, as described elsewhere (9). Special care was taken to cultivate Mφs in normoglycemic (5.5 mmol/l glucose) conditions in Dulbecco’s modified Eagle’s medium culture and to carefully monitor medium pH and lactate accumulation. For peritoneal Mφs, thioglycolate-elicited Mφs were prepared as described previously (23). The isolation of cardiomyocytes and cardiac fibroblasts from adult hearts was performed as previously described (24). In brief, 6- to 8-week-old B6 mice were injected intra-peritoneally with 0.5 ml heparin solution before being anesthetized with tribromoethanol. The excised heart was immediately mounted to a modified Langendorff apparatus and perfused with perfusion buffer, digestion buffer, and stopping buffer in sequential order. After enzyme dissociation, CMs were enriched through a gradient of calcium reintroduction buffer, and were plated accordingly for downstream experimentation.

**IN VITRO APOPTOSIS.** To compare relative effrocytosis efficiency across cell types, a common apoptotic stimulus was selected: staurosporine. CMs, cardiac fibroblasts, and Mφs were incubated with 0.5 μmol/l staurosporine. The morphology of apoptotic blebbing was quantified and confirmed by annexin V staining. Staurosporine was used as a robust means to induce apoptosis across different cell types (25) and also owing to our empirical observations that CMs are relatively resistant to ultraviolet light-induced apoptosis. Fibroblasts, primary Mφs, and CMs were induced to undergo apoptosis with 0.5 μmol/l of staurosporine treatment for 2 h followed by harvest of nonadherent floating cells (approximately 70% of the total cells). Apoptosis was confirmed by microscopic observation of cell size, morphology, and flow cytometric analysis of phosphatidylserine exposure by binding of annexin V-FITC (catalog no. 556491, BD Pharmingen, Franklin Lakes, New Jersey) according to the recommendations of the manufacturer (14).

**IN VITRO EFFEROCTYSIS ASSAY.** Peritoneal Mφs were plated at 2 × 10^5 in 24-well plates 1 day before the experiment. Apoptotic cells were prelabeled with Calcein AM before apoptosis was induced. The Mφ to CM ratio was 1:5. Cells were cocultured for 1 hour before 3 subsequent rinses to remove nonengulfed cells. The phagocytes were then fixed with 4% paraformaldehyde and analyzed by fluorescence microscopy.

**STATISTICAL ANALYSIS.** Data are presented as mean ± SEM Data fit a normal distribution. Thus, the Student t test was used to determine statistical significance and p < 0.05 were considered significant. SPSS Statistics (SPSS, Inc., Chicago, Illinois) version 17 was used for statistical analyses.

**RESULTS**

**EVIDENCE FOR CM-SPECIFIC CD47 EXPRESSION IN HUMANS AND CD47-INDUCTION AFTER EXPERIMENTAL MI.** A previous study characterized CD47 levels from total ventricular biopsy extracts (26);
however, our understanding of CM-specific expression in humans is limited. We, therefore, first used immunohistochemistry of human hearts after autopsy to test for CM-specific CD47. As shown in Figure 1A, human (Supplemental Table 1, Supplemental Figures 1 and 2) myocardial CD47 expression was found to colocalize to CMs; reduced colocalization was identified on other human cardiac cells, such as endothelial cells (Supplemental Figure 3). In specimens from patients who had succumbed to MI, CD47 staining intensity was interestingly increased (Figure 1B). Some of this CD47 colocalized with phagocytic ligand calreticulin (Supplemental Figure 4), leading us to hypothesize that increased CD47 might interfere with phagocytic efficiency (10). Indeed, increased human CM CD47 levels after acute ischemia is consistent with an impediment to dying myocyte phagocytic clearance and healing. Therefore, we set forth to measure CM CD47 under experimental ischemia and test if CD47 blockade could enhance myocyte phagocytosis and cardiac repair. Similar findings were seen in experimental mouse hearts, where the CD47 signal colocalized with CMs, in contrast with vimentin-positive cardiac fibroblasts (Figures 1C and 1D). To measure CM CD47 levels after MI, experimental mice were subjected to coronary occlusion, as described in the Methods. Interestingly, CD47 levels after MI were significantly increased in the ischemic area at risk versus the remote myocardium (Figure 1E). Ex vivo analysis corroborated these findings, revealing significant CD47 protein in isolated adult CMs. Interestingly, CD47 protein was further induced after treatment with stimuli that induced CM death (Supplemental Figure 5). Cell death-induced CD47 was confirmed on the cell surface, and in contrast with primary cardiac fibroblasts.
CD47-phagocytic regulation. To test the potential of CD47 blockade for rescuing basal CM phagocytosis inefficiency, we added anti-CD47 blocking antibodies to cocultures of primary adult differentiated murine CMs and Mφs. Figure 2A shows that CD47 blockade significantly heightened CM effecrocity, over and above isotype control, and specifically for apoptotic CMs (Figure 2B, Supplemental Figure 6). Effecroctic enhancements required CD47, because CMs prepared from Cdah7−/− mice were resistant to the effect of anti-CD47 blocking and Cdah7−/− CMs were more susceptible to phagocytic clearance at baseline (Figure 2C). These findings predicted similar responses by blocking CD47 ligand, Mφ SIRP-1z. Indeed, anti-SIRP-1z monoclonal antibody, added to Mφs, also increased phagocytosis (Figure 2D) of Cdah7−/− CMs. Similar findings were found when inducing induced pluripotent stem cell derived cardiomyocytes to apoptosis and cocultivating with human Mφs (Supplemental Figure 7).

Having tested proof of principle ex vivo, we next set out to test physiological relevance. Because of the potential therapeutic implications of our approach, we decided on a reperfusion model of acute murine MI. Also, because of potential nonphagocytic pathways impacted by CD47 signaling, we further decided to limit the blocking of CD47 to acute treatment. Thus, germline deficiency was neither a favorable strategy for our questions nor a viable approach, owing to previously published studies that indicated significant compensatory responses in Cdah7−/− mice (27).

**EX VIVO UPTAKE OF CM APOPTOTIC BODIES BY MφS IS REGULATED BY THE CD47–SIRPα AXIS.** We previously reported that phagocytosis is a significant early regulator of cardiac repair after an MI (9), and furthermore that CMs are engulfed by Mφs at a lower efficiency than other cell types that turnover after an MI (18). In that the Henson (10) and Weissman groups (11) have shown the potential of blocking CD47 to enhance phagocytosis, we reasoned that increased CD47 might render CMs particularly sensitive to

**IN VIVO TESTING.** Our experimental outline is shown in Supplemental Figure 8. Experimental B6 mice were subjected to ligation of the left anterior descending artery, followed by reperfusion, with or without anti-CD47 treatment. After performing a titration of anti-CD47 concentrations to find the optimal dose for affecting infarct size, we settled on 100 μg. In consideration of potential CD47–thrombospondin interactions, no significant changes in markers of neovascularization were measured (Supplemental Figure 9B). Furthermore, given that global Cdah7−/− knockout mice have been reported to exhibit an increase in arterial diastolic and systolic pressure and that CD47 has also been shown to suppress nitric oxide (NO) signaling in vascular cells after long-term thrombospondin binding (28), we assessed hemodynamics. However, no differences in systolic blood pressure were found after CD47 blockade (Supplemental Figure 9A).

**ACUTE CD47 BLOCKADE DURING REPERFUSION AFTER MI ENHANCES MYOCARDIAL PHAGOCYTOSIS.** To specifically measure effects on myocardial phagocytosis, we used mice transgenic for Mhc6-mCherry,
the fluorescence of which is restricted to CMs (18). In line with our hypothesis, mice administered anti-CD47 blocking antibody during reperfusion (Figure 3A), exhibited an increased profile of myocardial CD11b+ phagocytes with associated mCherry fluorescence (Figure 3A and 3B), an indicator of CM phagocytosis. Strikingly, mice administered adenoviral-associated virus CD47, which increased CD47 levels, exhibited reduced markers of myocardial phagocytosis (Figure 3C). Overexpression of CD47 did not affect infarct size, potentially owing to phagocytosis-independent signaling (Supplemental Figure 10). Importantly, anti-CD47 blocking strategies also led to other predicted consequences of enhanced clearance, namely, faster kinetics of cardiac innate immune cell resolution (Figure 3D), such as reduced Ly6cHI...
monocytes and heightened Ly6CLO monocytes cells (29). Because no differences in neutrophil turnover were found, this suggested specificity for CM uptake. Importantly, administration of antibodies did not affect initial levels or recruitment of blood-borne monocytes (Supplemental Figure 11).

THE EFFECTS OF ACUTE ANTI-CD47 ON INFARCT SIZE, SYSTOLIC FUNCTION, AND CARDIAC SCARRING. Impressively, singular administration of anti-CD47 suppressed infarct size and cardiac troponin release (Figures 4A to 4C). Blockade of CD47 also enhanced systolic cardiac function as indicated by increased LV ejection fraction (Figures 5A to 5C). Anti-CD47 monoclonal antibody treatment limited ventricular remodeling, because treated mice exhibited reduced scarring, demonstrated by a 2-fold reduction in collagen fraction after Masson’s Trichrome staining (Figure 5D). Taken together, our data suggest a significant potential for incorporating CD47 blocking strategies toward the amelioration of cardiac injury. This approach is unique in its association and targeting of phagocytosis-associated repair.

DISCUSSION

Although cardiac reperfusion therapies have reduced patient mortality after AMI, improved survival has also led to increases in the frequency of heart failure. A contributing factor to both the benefit and detriment of cardiac repair after ischemia and reperfusion is the innate immune response. Immune cells directly interact with the injured tissue and this includes through binding to the molecular ligand CD47 on parenchymal cells. Herein, we target CD47 with blocking antibodies to enhance phagocytic clearance of dying CMs and in turn, augment cardiac wound healing after MI. Although the gold standard to test molecular requirements is the genetic knockout, CD47 germline deficiency tolerizes mice to the CD47-Sirpα-phagocytosis pathway, rendering phagocytosis CD47 independent (27). In this context, CD47-blocking strategies have most notably been pioneered by the Weissman group and to promote tumor cell uptake (11). During wound healing, CD47 inhibition accelerates wound closure after dermal thermal injury in mice, independent of CD47 ligand thrombospondin-1 (30). Herein, a similar approach is tested in the heart, but with distinct mechanistic and clinical implications. Although multiple cell types turnover after MI (31–33), distinct myocardial patterns of expression during ischemia (Figure 1, Supplemental Figure 4) and CM-specific mechanisms may render CMs uniquely sensitive to the effects of CD47-blockade. Also, dying cells must both downregulate anti-phagocytic signals such as CD47 and concomitantly induce pro-phagocytic signals. Thus, strategies that target CD47 for phagocytosis should be specific to dying cells, which further require induction of phagocytic signals.

Previous studies suggest that phagocytosis of adult CMs by Mφs is naturally inefficient (18). Indeed, diseases of aging such as MI do not exert evolutionary pressure to optimize CM clearance after severe MI. During aging, phagocytosis is compromised (34), including in the heart (35). In the case of CMs, we speculate that low phagocytic efficiency is linked to the low regenerative potential of adult myocytes and could require enhanced mechanisms to prevent phagocytic removal. For example, our data indicate that CD47 surface levels are uniquely heightened after apoptosis (Supplemental Figure 5) and our unpublished experiments suggest novel interactions with membrane scaffolding proteins, which may in turn regulate CD47 cell surface half-life and interactions with other phagocytic molecules, including calreticulin (Supplemental Figure 4).
CM, the function, spatial organization, and interactome of CD47 remains understudied.

Phagocytosis-targeting approaches might be enhanced by the administration of osonins. For example, natural defects in efferocytosis in cardiovascular disease have been hypothesized to be the result of deficient Gas6 expression (36), a critical bridging molecule that facilitates Mφ-mediated engulfment. Similarly, a polymorphism associated with coronary artery disease leads to reduced calreticulin expression on vascular smooth muscle cells and decreased efferocytosis (37). On the Mφ side, blocking SIRP-1ζ may also enhance phagocytic efficiency. This approach is attractive because of the potential to target peripheral blood phagocytes, destined for the heart. Recent advances include the engineering of high-affinity SIRPz variants, which are capable of enhancing CD47 blockade (38). However, other studies indicate SIRP-1ζ is critical to platelet activation (39) and neutrophil and Mφ migration (40), potentially precluding their efficacy in this regard. Furthermore, cardiac SIRP-1ζ protects against myocardial hypertrophy through disruption of TLR4 signaling (41).

**STUDY LIMITATIONS.** This study has several limitations, including the restricted extent to which murine studies can be extrapolated to humans. Further studies are necessary to test CD47 blockade in large animal...
models and examine conservation and reproducibility across species. Also, a closer look at our echocardiography data reveals that the slope of the reduction in the ejection fraction in the anti-CD47 cohort was greater than that in the control group (Figure 5). Therefore, additional studies should test the chronic benefit versus the transient nature of this treatment. It is important to consider phagocytosis-independent effects of CD47 blockade. For instance, CD47 targeting during tissue ischemia and reperfusion injury is associated with angiogenesis (42) and effecrocytosis in heart can trigger vascular endothelial growth factor A (43). However, we did not detect significant increases in vascular density (Supplemental Figure 9). CD47 also has an inhibitory effect on NO signaling. Hypoxia can induce NO synthases and increased NO signaling can improve cell survival under ischemia, in CD47-null mice. In endothelial cells, NO production induces stimulation of vascular smooth muscle cells to promote vascular relaxation; however, induced CD47 ligand thrombospondin-1 inhibits NO production and stimulates production of reactive oxygen species (44). In our own hands, CD47 blockade does not affect CM viability in vitro. Also, the effects on blood pressure were not found during CD47 blockade (Supplemental Figure 9). Finally, excessive phagocytic uptake has been shown to be detrimental in the central nervous system through phagocytosis-induced cell death (45). In this context, CD47 may in some instances act as a prophagocytic signal (46).

CONCLUSIONS

An increased understanding of the basic mechanisms of CD47 regulation in CMs holds the potential to unenroll nonphagocytic CD47 signaling to selectively enhance the clearance of dying cells after MI and improve tissue repair. The acute nature of infarct-associated CM death, paired with standards of percutaneous coronary intervention, offer a tractable opportunity for CD47-targeted approaches during clinical reperfusion.

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PERSPECTIVES

COMPETENCY IN MEDICAL KNOWLEDGE: CD47, also known as integrin-associated protein, is known to regulate multiple distinct cellular processes, including the inhibition of phagocytosis of dying cells, or effecrocytosis, by immune cell Mφs. Our preclinical studies indicate that CD47-blocking antibodies enhance the phagocytic removal of dying cardiac myocytes, after reperfusion in MI. This intervention also improved the resolution of cardiac inflammation and reduced infarct size.

TRANSLATIONAL OUTLOOK: Further studies are needed to enhance the pharmacological route of administration for anti-CD47 inhibitors and to test the efficacy of humanized anti-CD47 antibodies.

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**KEY WORDS** phagocytosis, CD47, macrophage

**APPENDIX** For a supplemental table and figures, please see the online version of this article.