An Injectable Oxygen Release System to Augment Cell Survival and Promote Cardiac Repair Following Myocardial Infarction

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Oxygen deficiency after myocardial infarction (MI) leads to massive cardiac cell death. Protection of cardiac cells and promotion of cardiac repair are key therapeutic goals. These goals may be achieved by re-introducing oxygen into the infarcted area. Yet current systemic oxygen delivery approaches cannot efficiently diffuse oxygen into the infarcted area that has extremely low blood flow. In this work, we developed a new oxygen delivery system that can be delivered specifically to the infarcted tissue, and continuously release oxygen to protect the cardiac cells. The system was based on a thermosensitive, injectable and fast gelation hydrogel, and oxygen releasing microspheres. The fast gelation hydrogel was used to increase microsphere retention in the heart tissue. The system was able to continuously release oxygen for 4 weeks. The released oxygen significantly increased survival of cardiac cells under the hypoxic condition (1% O₂) mimicking that of the infarcted hearts. It also reduced myofibroblast formation under hypoxic condition (1% O₂). After implanting into infarcted hearts for 4 weeks, the released oxygen significantly augmented cell survival, decreased macrophage density, reduced collagen deposition and myofibroblast density, and stimulated tissue angiogenesis, leading to a significant increase in cardiac function.

MI causes massive death of cardiac cells including cardiomyocytes, cardiac fibroblasts and endothelial cells. Extremely low oxygen content in the infarcted area is a major cause of death. MI also induces severe pathogenic inflammatory responses, scar formation, and cardiac function decrease. Protection of cardiac cells and promotion of cardiac repair are key treatment goals. These goals may be achieved by clinical reperfusion intervention that reintroduces oxygen into the infarcted heart. However, not all patients are eligible for this type of intervention. Cell therapy has potential to use endogenous or exogenous cells for cardiac repair, yet cell survival is inferior in the low oxygen condition of the damaged hearts. Biomaterial therapy with or without growth factors may aid myocardial repair by providing mechanical support to the heart tissue, and affecting tissue inflammation and angiogenesis. However, the efficacy remains low due to their inability to provide oxygen to metabolic-demanding cardiac cells at early stage of tissue damage. To address the critical need of oxygen to protect cardiac cells, direct supply of sufficient oxygen in the infarcted area while not provoking deleterious effects is necessary. However, this cannot be achieved by current oxygen therapy approaches.

Oxygen supplementation is a standard treatment for MI patients because it increases oxygen level in the blood of healthy tissues to avoid hypoxic damage caused by lower blood pumping ability after MI. It may also augment oxygen level in the infarcted tissue to protect cardiac cells although this area has extremely low blood supply. As a result, cardiac function may improve. Experiments using canine model have demonstrated that inhalation of 100% oxygen decreased infarct size and increased cardiac function (ejection fraction). Several clinical studies also showed similar effects when patients inhaled 100% oxygen, yet some did not show any effect.

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Hyperbaric oxygen therapy uses 100% oxygen with high pressure (>1 atm). The purpose is to better increase blood oxygen level than traditional oxygen therapy\(^{25-27}\). Animal studies have shown that hyperbaric oxygen therapy increased cell survival in the infarcted hearts\(^{26,37}\). Some clinical studies demonstrated that hyperbaric oxygen therapy decreased end-systolic volume by 20% and increased cardiac output by 10%\(^{38}\). Yet other clinical studies did not have similar beneficial effects\(^{39,40}\). Intracoronary injection of arterial blood supersaturated with oxygen is also an approach to augment oxygen level in the infarcted area. Some clinical studies demonstrated that this approach can significantly improve cardiac function after 30 days for patients with large damaged area\(^{41-43}\). However, no positive effect was found in some other clinical studies\(^{41-43}\). Transfusion of oxygen carriers into blood after MI to increase blood oxygen level has been tested in animal models. The results demonstrated that infarct size was reduced and cardiomyocyte survival was increased\(^{44-47}\). However, clinical data on this approach is lacking.

Overall, current oxygen therapy for MI treatment is focused on systemic oxygen delivery, and the therapeutic efficacy is low. In addition, the results are inconsistent in clinical trials and preclinical studies\(^{27-29}\). This is because: (1) The infarcted area has extremely low blood flow, thus largely limiting oxygen in the blood to diffuse into the area\(^{48}\). The oxygen level may be too low to protect substantial number of cells; (2) systemic increase of blood oxygen level decreases coronary artery blood flow\(^{49,50}\). This directly decreases oxygen diffusion to the infarcted area; and (3) current approaches cannot increase oxygen level in the blood for prolonged period to continuously provide oxygen to the cardiac cells since the oxygen level decreases to the normal level shortly after the treatment. Prolonged inhalation of oxygen or injection of oxygen carriers into blood may lead to side effects on healthy tissues as the oxidative stress may be increased in these tissues causing cell death and tissue inflammation.

To address limitations of current oxygen therapy in order to largely augment its therapeutic efficacy, an approach that can efficiently deliver necessary level of oxygen to the infarcted area for a prolonged period while not causing side effects is critically necessary. In this work, we developed a new oxygen delivery system that can be delivered specifically to the infarcted tissue in order to avoid side effects on healthy tissues, and can gradually release sufficient oxygen to continuously oxygenate the metabolic-demanding cardiac cells. The effect of released oxygen on cardiac cell survival, myofibroblast formation, cardiac fibrosis, tissue inflammation, angiogenesis and cardiac function was studied.

**Results**

**Fabrication of hydrogel-based oxygen release system and oxygen release kinetics.** The oxygen release system was based on a thermal sensitive and fast gelation hydrogel, and oxygen release microspheres. The hydrogel polymer had NIPAAm/HEMA/AOLA ratio of 86.0/10.5/3.5 as determined from \(^1\)H-NMR spectrum. The 20% hydrogel solution was flowable at 4 °C, and can be readily injected through a 28-gauge needle. The hydrogel solution exhibited a thermal transition temperature of 26.7 °C. The gelation time was less than 7 s at 37 °C. After incubation in the DPBS for 4 weeks at 37 °C, the hydrogel showed 5.4% of weight loss.

The oxygen release microspheres were fabricated by electrospaying (Fig. 1), and had core-shell structure with PLGA as shell and PVP/H\(_2\)O\(_2\) complex as core (Fig. 2). Oxygen was generated after the PVP/H\(_2\)O\(_2\) complex released from microspheres was converted into molecular oxygen and water by catalase encapsulated in the hydrogel. The released oxygen was measured by using oxygen-sensitive luminophore Ru(PH\(_{2}\)phen\(_3\))Cl\(_2\) and oxygen-insensitive fluorophore rhodamine-B. The Ru(PH\(_{2}\)phen\(_3\))Cl\(_2\) has high quantum yield and oxygen quenching, and is thermal stable at physiological temperature\(^{51,52}\), while rhodamine-B allows for correction of typical errors, such as heterogeneous fluorophore concentration, bleaching, and fluctuations in excitation light intensity\(^{53}\). Figure 2 demonstrates that the oxygen release system was able to continuously release oxygen in a 4-week testing period. The release showed a two phase pattern. In the first 7 days, the oxygen level was gradually increased. After 1 day of release, the oxygen level was above 5%. The release was peaked at day 7 where oxygen level was 24.8%. After 7 days, the oxygen level was steadily decreased. By day 28, it was 11.2%.

**Effect of oxygen release on cell survival under hypoxic conditions.** To evaluate the survival of cardiomyocytes, endothelial cells and cardiac fibroblasts under hypoxic condition, each cell type was first seeded into 3D collagen gels. The oxygen release system was then injected into the collagen gels followed by culturing under 1% \(O_2\) condition (Gel/O\(_2\) Hypoxia group). Controls were collagen gels injected with hydrogel only and cultured under hypoxia (Gel Hypoxia group) or normal oxygen condition (Gel Normal group). Figure 3A shows that 1% \(O_2\) condition significantly decreased cell dsDNA content after 2 days of culture (\(p < 0.0001\) for cardiomyocytes, and \(p < 0.01\) for cardiac fibroblasts and HUVECs Fig 3A), demonstrating that this oxygen condition caused cell death. The injection of hydrogel only did not increase dsDNA content for 3 cell types (\(p > 0.05\)). In contrast, the injection of oxygen release system significantly augmented cell dsDNA content (\(p < 0.001\) in cardiomyocytes, \(p < 0.01\) in HUVECs, and \(p < 0.05\) for cardiac fibroblasts), suggesting that the released oxygen increased cell survival under extremely low oxygen condition. The dsDNA content for each cell type was similar to that cultured under normal oxygen condition (Fig. 3A). To observe live cells, they were stained with a live cell tracker CMFDA before seeding into collagen gels. Consistent with dsDNA data, the cells cultured under 1% \(O_2\) condition with the injection of hydrogel showed dramatic lower cell density than those cultured under normal oxygen condition (Fig. 3B). The injection of oxygen release system increased cell density (Fig. 3B).

**Effect of oxygen release on cell ROS content.** To determine whether oxygen release affected cell ROS content, cardiomyocytes, HUVECs and cardiac fibroblasts were stained with a ROS sensitive dye CM-H\(_2\)DCFDA before seeding into collagen gels. The relative ROS content was quantified. The 3 cell types showed different ROS contents under 3 different conditions (Fig. 4). At low oxygen condition (1% \(O_2\)), the ROS contents were significantly decreased for each cell type (\(p < 0.0001\)) compared to the normal oxygen condition. Injection of the oxygen
release system into the collagen gels significantly increased cell ROS content ($p < 0.0001$). However, the ROS content was similar to that of cells cultured under normal culture conditions ($p > 0.1$).

**Effect of oxygen release on myofibroblast formation under hypoxic condition.** To determine the effect of oxygen release on myofibroblast formation, the cardiac fibroblasts were seeded in the 3D collagen gel that can maintain fibroblast phenotype for at least 48 h, and cultured in the medium supplemented with TGF-β1 under either normal oxygen or 1% O$_2$ condition. After 2 days of culture, the expressions of TGF-β1 and TGF-βRII at the mRNA level were significantly upregulated in the 1% O$_2$ condition than in the normal oxygen condition ($p < 0.05$ for both expressions Fig. 5). This increases the possibility of TGF-β1 binding to TGF-βRII, leading to the formation of myofibroblasts. Injection of oxygen release system into the collagen gel significantly downregulated the expressions of these genes ($p < 0.05$ Fig. 5).

To characterize cardiac fibroblast differentiation into myofibroblast, myofibroblast markers including αSMA, CTGF, and collagen1A1 were quantified at the mRNA level. The 3 markers were significantly downregulated after injection of oxygen release system into the collagen gel (Gel vs. Gel/O$_2$ groups. $p < 0.05$ Fig. 6A). At the protein level, nearly all of the cells were αSMA positive myofibroblasts in the Gel group (Fig. 6B and D). The release of oxygen in the collagen gel (Gel/O$_2$ group) significantly decreased the ratio of myofibroblasts to 49% ($p < 0.05$ Fig. 6C and D). The increase of myofibroblast ratio is not caused by gel as we found that cardiac fibroblasts cultured on the gel surface without adding TGF-β1 did not express αSMA (data not shown).

**Effect of oxygen release system on infarcted left ventricular wall thickness after MI.** To examine the effect of released oxygen on the remodeling of infarcted left ventricles, hydrogel encapsulated with oxygen release microspheres and catalase (Gel/O$_2$ group) was injected into the infarcted rat hearts 30 min after MI. Sham, infarcted hearts without treatment (MI group), and infarcted hearts injected with hydrogel only (Gel group) were used as controls. Four weeks after injection, substantial left ventricular dilation and wall thinning were found for the MI group (Fig. 7). Injection of Gel and Gel/O$_2$ groups appeared to have less dilation and wall thinning (Fig. 7). The thickness of myocardium at infarcted area was quantified from H&E images, and normalized to the left ventricle thickness of the Sham group. Figure 7 demonstrates that wall thickness of the MI group was only 30.2% of the Sham group at 4 weeks post MI. Injection of hydrogel only (Gel group) significantly increased wall thickness.
thickness to 47.4% of the Sham group (p < 0.001, Gel group vs. MI group). Injection of oxygen release system further increased wall thickness to 58.0% of the Sham group (p < 0.01, Gel/O₂ group vs. Gel group). The above results suggested that injection of hydrogel and oxygen release system attenuated adverse remodeling after MI.

In addition, higher magnification H&E images of the infarcted area demonstrated that cells infiltrated into the hydrogel (Fig. 7).

**Effect of oxygen release system on host cell survival and proliferation.** Cell survival was quantified for cardiomyocytes in the infarcted area at 4 weeks post MI. The surviving cardiomyocytes were identified as MHC positive cells (Fig. 8). The lowest cell density was found for the MI group where only few rounded cardiomyocytes were observed. The injection of hydrogel significantly increased cardiomyocyte density in the infarcted area (p < 0.05). Most of cells assumed round morphology with small number of cells was elongated. The injection of oxygen release system largely increased cardiomyocyte survival as the cell density was significantly greater than the Gel group (p < 0.01, Gel/O₂ group vs. Gel group). Interestingly, most of the surviving cardiomyocytes were elongated with similar morphology as those in the Sham group.

**Figure 2.** Core-shell structure of oxygen release microsphere and oxygen release kinetics. (A) PLGA shell; (B) PVP/H₂O₂ core; (C) combined core-shell structure. For imaging purpose, rhodamine and draq5 were added to the shell and core respectively before fabrication; (D) SEM image of microsphere; and (E) oxygen release kinetics during a 28-day release period. Scale bar = 5 µm.
To quantify host cell proliferation, Ki67 staining on the infarcted area was performed and Ki67+ cells were quantified (Fig. 9). After 4 weeks of injection, the MI group showed the minimum cell proliferation. Injection of hydrogel substantially increased cell proliferation ($p > 0.05$, Gel group vs. MI group). The most significant cell proliferation was observed for the Gel/O$_2$ group that had oxygen release ($p < 0.01$, Gel/O$_2$ group vs. Gel group).

**Figure 3.** HUVECs, cardiac fibroblasts, and cardiomyocytes survival under normal and hypoxic conditions. (A) dsDNA content. Three cell types were seeded in 3D collagen gel, respectively. Hydrogel (Gel) was then injected into the collagen gel and cultured under normal (Gel Normal group) or 1% O$_2$ (Gel Hypoxia group) conditions. Oxygen release system (Gel/O$_2$) was also injected into the collagen gel but cultured under 1% O$_2$ (Gel/O$_2$ Hypoxia group). dsDNA was measured 2 days after injection. dsDNA under hypoxia was normalized to the collagen gel injected with Gel and cultured under normal condition; and (B) live cell images of HUVECs, cardiac fibroblasts, and cardiomyocytes in the 3D collagen gel. Cells were labeled with live cell tracker CMFDA before seeding. $^*p < 0.05$. Scale bar = 30 µm.
Effect of oxygen release system on cardiac fibrosis. Cardiac fibrosis is characterized by increased myofibroblast density and collagen content in the infarcted tissue. Myofibroblasts are mainly responsible for collagen secretion. To determine the efficacy of oxygen release on myofibroblast formation, infarcted tissue was stained for αSMA and vWF (Fig. 10). Myofibroblasts are αSMA+ cells that are not co-localized with vWF+ endothelial cells in the vessels. Figure 10 demonstrates that myofibroblast density was significantly lower in the Gel/O₂ group than in the MI and Gel groups (p < 0.01, Gel/O₂ group vs. Gel group or MI group). The injection
of hydrogel did not affect myofibroblast formation compared to MI group (p > 0.1). Picrosirius red staining was utilized to identify collagen in the infarcted area (Fig. 11). Total collagen content in the MI and Gel groups was significantly higher than that in the Gel/O₂ group (p < 0.05, Gel/O₂ group vs. MI and Gel groups). There was no significant difference between MI and Gel groups (p > 0.05). The above results demonstrated that oxygen release attenuated cardiac fibrosis.

Effect of oxygen release system on tissue inflammatory and ROS content in infarcted hearts.

Tissue inflammation in the infarcted area was characterized at 4 weeks post MI by F4/80 positive macrophages (Fig. 12). The percentage of F4/80 positive cells was similar in MI group and Gel group. Injection of oxygen release system into the infarcted hearts significantly decreased percentage of F4/80 positive cells compared to MI and Gel groups (p < 0.01, Gel/O₂ group vs. MI and Gel groups).

To determine whether oxygen release in the infarcted hearts increased ROS content in the tissue, ROS staining was performed in the MI, Gel, and Gel/O₂ groups (Fig. 13). The mean gray value of each cell in each surgery group was measured to evaluate the total ROS content in each group. MI group exhibited the highest ROS content. Injection of hydrogel slightly decreased ROS content (p > 0.5, Gel group vs. MI group). Interestingly, the injection of oxygen release system did not increase but significantly decreased ROS content (p < 0.01). The above results demonstrated that oxygen release in the infarcted heart tissue significantly attenuated tissue inflammation and ROS formation.

**Figure 5.** Gene expression of TGFβ1 (A) and TGFβRII (B) in cardiac fibroblasts when seeded in collagen gels. The Gel and Gel/O₂ groups were injected into the collagen gels respectively and cultured under normal condition or hypoxia (1% O₂). Gene expression was measured 2 days after injection. TGFβ1 (5 ng/mL) was added into the medium during culture. *p < 0.05.
Effect of oxygen release on infarcted tissue angiogenesis and cardiac function. To investigate if oxygen release can increase infarcted tissue angiogenesis, vessel density was quantified for the MI, Gel, and Gel/O2 groups (Fig. 10E). The MI and Gel group had similar vessel density (p > 0.1). The injection of oxygen release system remarkably increased vessel density compared to MI and Gel groups (p < 0.01, Gel/O2 group vs. Gel and MI groups). Cardiac function was characterized by echocardiography. Ejection fraction (EF), fractional shortening (FS), and fractional area change (FAC) were quantified (Fig. 14) 4 weeks of injection, the Gel group exhibited substantially greater EF, FS, and FAC than the MI group (p > 0.05). Compared to the Gel group, the Gel/O2 group showed significantly higher EF, FS, and FAC (p < 0.05 for EF, FS and FAC). These results suggested that oxygen release promoted tissue angiogenesis and increased cardiac function.

Discussion
In this work, an oxygen release system was developed to rescue cardiac cells and promote cardiac repair after acute MI. The system was able to sustained release oxygen to continuously oxygenate the metabolic-demanding cardiac cells and augment their survival. Controlled oxygen release to heart tissue cannot be achieved by current oxygen delivery approaches like hyperbaric oxygen therapy where tissue oxygen content decreases shortly after the treatment is finished. The oxygen delivery system was specifically injected into the infarcted tissue. The localized oxygen delivery is advantageous over systemic oxygen delivery as higher tissue oxygen content may be achieved. Systemic oxygen delivery relies on blood flow to transport oxygen to the infarcted hearts. Yet the damaged heart tissue is poorly vascularized.

The oxygen release system was based on a thermosensitive and fast gelation PNIPAAm hydrogel, and oxygen release microspheres. The hydrogel was served as carrier for the oxygen release microspheres. PNIPAAm based...
Hydrogels have good biocompatibility and can solidify at 37 °C without using crosslinkers.\textsuperscript{55,56} The hydrogel solution in this work was able to solidify within 7 s at 37 °C. This is beneficial for myocardial injection as the hydrogel may quickly immobilize in the heart tissue, thus increasing the retention of encapsulated microspheres in the tissue. This is evidenced in the in vivo study where hydrogel leaking was not observed during myocardial injection. The hydrogel solution had a thermal transition temperature of 26.7 °C and can be readily injected through 28-gauge needle. The hydrogel was degradable. The degradation product had gelation temperature of 41.2 °C.

Figure 7. H&E staining and relative wall thickness of the infarcted hearts 4 weeks after injection. (A–D) H&E staining of Sham (A); MI (B); Gel (C); and Gel/O\textsubscript{2} (D) groups. Scale bar = 2 mm for whole heart view, and scale bar = 50 µm for higher magnification view. The higher magnification H&E images indicated that cells were infiltrated into the hydrogel; and (E) relative wall thickness. $p < 0.05$. 

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can thus dissolve in the body fluid and be removed from body. These properties allow the developed hydrogel to be suitable for cardiac therapy.

The oxygen release microspheres assumed core-shell structure with PVP/H$_2$O$_2$ as core and PLGA as shell (Fig. 2). The mechanism of oxygen release is that the PVP/H$_2$O$_2$ complex diffuses through the PLGA shell, and the H$_2$O$_2$ is converted into molecular oxygen by catalase in the hydrogel (Fig. 1). The oxygen was able to continuously release from the hydrogel for 4 weeks (Fig. 2). After 1 day of release, the oxygen level was reached 5%. The peak release was observed on day 7 where the oxygen level was 24.8%. Oxygen release may be faster in vivo as the

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**Figure 8.** MHC staining and MHC$^+$ cell density of the infarcted hearts 4 weeks after injection. (A–D) MHC staining of Sham (A); MI (B); Gel (C); and Gel/O$_2$ (D) groups; and (E) MHC$^+$ cell density in the infarcted area. Sham, MI and Gel groups were used as controls. Scale bar = 30µm. *p < 0.05.
enzymes in the infarcted hearts may accelerate the degradation of hydrogel and PLGA. However, it is possible that oxygen release can last for 4 weeks since the hydrogel was observed after 4 weeks of implantation.

This oxygen release system has advantages over those MgO$_2$, CaO$_2$, H$_2$O$_2$ (without forming complex), and fluorinated compounds based systems. First, the duration of release is longer. Oxygen release in the above mentioned systems typically lasted for 2 weeks or shorter. In contrast, oxygen release in our system persisted for 4 weeks. Second, our oxygen release system is safer as the released H$_2$O$_2$ can be quickly converted into oxygen by catalase. Therefore no free H$_2$O$_2$ is released from the system. Free H$_2$O$_2$ may be toxic to cells. The MgO$_2$, CaO$_2$, and H$_2$O$_2$ (without forming complex) based oxygen release systems release free H$_2$O$_2$, and rely only on H$_2$O$_2$ decomposition to form oxygen. Third, our system is more suitable for cardiac therapy than the MgO$_2$ and CaO$_2$-based oxygen release systems since it does not generate side product Mg$^{2+}$ or Ca$^{2+}$, which may lead to an abnormal ion transient in the heart tissue.

Various studies have shown that a relatively low oxygen condition (5–20%) may not affect cell survival, however, the extremely low oxygen condition (<1%) in infarcted hearts leads to extensive cell death. Salvation of major cardiac cells like cardiomyocytes, endothelial cells and cardiac fibroblasts is critical for cardiac repair after MI. To investigate the effect of oxygen release on cardiac cell survival under low oxygen condition mimicking that of the infarcted hearts, cells were seeded into a 3D collagen model and cultured under 1% oxygen condition. This oxygen concentration has been commonly used to simulate the pathological hypoxic condition in infarcted hearts. Cardiomyocytes, cardiac fibroblasts and endothelial cells experienced significant death under 1% oxygen condition after 2 days of culture (Fig. 3). The injection of oxygen release system significantly augmented the survival of all 3 cell types (Fig. 3). The improved cell survival is likely due to cell oxygen content increase. One of the limitations of 3D collagen model is that cell survival can only be evaluated for 2 days due to substantial shrinkage.

Consistent with in vitro results, injection of the oxygen release system into infarcted hearts increased cardiomyocyte survival. Figure 8 demonstrates that the number of MHC positive cardiomyocytes was significantly greater in the group with oxygen release (Gel/O$_2$ group) than that without oxygen release (Gel group). In addition,

Figure 9. Ki67 staining of the infarcted hearts harvested 4 weeks after injection. (A–C) Ki67 staining of MI (A); Gel (B); and Gel/O$_2$ (C) groups; and (D) Ki67 $^+$ cell density in the infarcted area. Scale bar = 30 µm. *p < 0.05.
The survived cardiomyocytes in the oxygen release group showed similar morphology as the healthy cardiomyocytes in the Sham group (Fig. 8). Oxygen release also promoted cell proliferation in the infarcted region (Fig. 9). Ki67 staining results demonstrated that the density of Ki67 positive cells in the Gel/O2 group was significantly higher than in Gel group. Since adult cardiomyocytes do not proliferate, the Ki67 positive cells likely contain endothelial cells and cardiac fibroblasts. This is indirectly evidenced by greater vessel density in the Gel/O2 group than in the Gel group (Fig. 14).

**Figure 10.** Immunohistological analysis of the infarcted region and myofibroblast density after 4 weeks of injection. (A–C) Immunohistological staining of αSMA (green), vWF (red), and Hoechst (blue) for MI (A); Gel (B); and Gel/O2 (C) groups; (D) myofibroblast density quantified from immunohistological images; and (E) blood vessel density in the infarcted area. Scale bar = 60 μm. *p < 0.05.
One of the concerns when using oxygen therapy is ROS generation as high oxygen concentration may increase ROS content in cells and tissues, leading to cell death. Within the statistical power of the \textit{in vitro} study in Fig. 4, ROS content of cardiomyocytes, cardiac fibroblasts, and endothelial cells in the oxygen release group did not significantly increase compared to that in the normal oxygen condition. Interestingly, the released oxygen significantly decreased ROS content in the infarcted hearts after 4 weeks of implantation (Fig. 13). These \textit{in vitro} and \textit{in vivo} results demonstrated that the released oxygen concentration was appropriate for myocardial repair. The reduced ROS content in the infarcted tissue is possibly resulted from decreased macrophage density (Fig. 12). Macrophages are main source of ROS after MI\textsuperscript{80}.

Decrease of myofibroblast formation is critical to reduce cardiac fibrosis. After MI, TGF\textbeta\textsuperscript{3} expression is upregulated. This promotes myofibroblast formation primarily through TGF\textbeta\textsuperscript{3} binding to TGF\textbeta\textsuperscript{3}RIIs on cardiac fibroblasts\textsuperscript{81}. In the infarcted hearts, cardiac fibroblast is one of the major sources of TGF\textbeta\textsuperscript{3}81. We found that the extremely low oxygen condition (1% O\textsubscript{2}) significantly upregulated the expressions of TGF\textbeta\textsuperscript{3} and TGF\textbeta\textsuperscript{3}RII on cardiac fibroblasts. However, oxygen release significantly downregulated both expressions (Fig. 5). This reduces the possibility of TGF\textbeta\textsuperscript{3} binding to TGF\textbeta\textsuperscript{3}RII of cardiac fibroblasts resulting in decreased myofibroblast density. It is confirmed by \textit{in vitro} cardiac fibroblast differentiation study where oxygen release significantly decreased \alpha\textsuperscript{SMA}, CTGF and collagen1A1 expressions at mRNA level, and \alpha\textsuperscript{SMA} expression at protein level (Fig. 6). After injecting into infarcted hearts for 4 weeks, the oxygen release group (Gel/O\textsubscript{2} group) showed significantly lower myofibroblast density and total collagen content than the Gel group that had no oxygen release (Figs 10 and 11). This is consistent with \textit{in vitro} findings. These results demonstrated that oxygen release significantly decreased myofibroblast formation and attenuated cardiac fibrosis after MI.

To examine the effect of oxygen release on left ventricular geometry after MI, the oxygen release system (Gel/\textsubscript{O\textsuperscript{2}} group) was injected into the infarcted area. MI and Gel groups were used as controls. Four weeks after injection, wall thickness of the Gel group was significantly greater than that of the MI group (Fig. 7). This is due to bulking effect of the hydrogel\textsuperscript{82}. Similar effect was reported in other studies using different types of hydrogels\textsuperscript{83,84}. The injection of Gel/O\textsubscript{2} group further increased wall thickness (Fig. 7). This increase is likely the contribution of hydrogel as well as cardiac cells saved by the released oxygen. Figures 8 and 9 demonstrate that densities of the surviving cardiomyocytes and proliferating cells were significantly higher in the Gel/O\textsubscript{2} group. It is also possible that the enhanced angiogenesis contributed to the wall thickness increase as the Gel/O\textsubscript{2} group had significantly
greater vessel density than the Gel group (Fig. 10E). The released oxygen may promote endothelial cell survival and proliferation under ischemic conditions of the infarcted hearts, resulting in the enhanced angiogenesis.

Cardiac function was examined 4 weeks post-MI. Injection of hydrogel only (Gel group) slightly increased EF, FS and FAC compared to the MI group. This is likely attributed to the wall stress decrease after hydrogel injection. Previous studies have shown that applying external matrix with suitable mechanical properties can effectively reduce the evaluated wall stress after MI84. The injection of oxygen release system significantly improved cardiac function (EF, FS, and FAC) compared to the Gel group (Fig. 14). The increase in cardiac function in the oxygen release group is possibly resulted from the mechanical support of hydrogel, increased cell survival and proliferation, greater angiogenesis, reduced inflammation, and decreased cardiac fibrosis. Overall, the above results demonstrated the benefits of using controlled oxygen release to protect cardiac cells and promote myocardial repair after MI.

The developed oxygen release system may be delivered into the infarcted hearts during coronary revascularization after MI85. Acute MI model was used in this work to evaluate the efficacy of oxygen release in promoting cardiac cell survival and cardiac repair. The major limitation of this model is its relatively low clinical relevance since patients may not be able to receive this type of treatment shortly after MI. However, this animal model has been widely used to investigate the efficacy of cell and biomaterial therapies after MI84–88.

Methods

Materials. All materials were purchased from Sigma-Aldrich unless otherwise stated. N-isopropylacrylamide (NIPAAm, TCI) was recrystallized in hexane before polymerization. 2-hydroxyethyl methacrylate (HEMA, TCI) was used after removing inhibitors through a column filled with inhibitor remover. Poly(lactide-co-glycolic acid) (PLGA) with LA/GA ratio of 50/50 ratio and inherent viscosity of 0.55–0.75 dL/g was purchased from Lactel. Hydrogen peroxide (30% aqueous solution), poly(N-vinylpyrrolidone) (PVP, Fisher Scientific) with molecular weight of 40–90 kDa, and bovine liver catalase (2000–5000 units/mg) were used as received.

Synthesis and characterization of hydrogel. The hydrogel was synthesized by free radical polymerization of NIPAAm, HEMA, and a macromer acrylate-oligolactide (AOLA). In brief, NIPAAm, HEMA, and AOLA with molar ratio of 86/10/4 were dissolved in 150 mL 1,4-dioxane under the protection of nitrogen. Initiator
benzoyl peroxide was then added. The polymerization reaction was conducted at 70 °C for 24 h with continuous stirring. The solution was precipitated in hexane. The obtained polymer was purified twice using THF as dissolving solvent and diethyl ether as precipitation solvent.

Hydrogel solution (20 wt%) was prepared by dissolving the synthesized polymer in Dulbecco's modified phosphate buffer saline (DPBS) at 4 °C. The thermal transition temperature of the solution was measured utilizing DSC over a temperature range of 0 °C and 60 °C with a heating rate of 10 °C/min. Injectability of the hydrogel solution was tested by injecting the solution through a 28-gauge needle typically used for tissue injection. Gelation time of the hydrogel solution was determined using a temperature controllable Olympus microscope. The time needed for the 4 °C hydrogel solution to become completely opaque at 37 °C was considered as gelation time. Hydrogel degradation was conducted in DPBS at 37 °C for 4 weeks. Weight loss was determined. At least 5 samples were used for degradation study.

**Fabrication of oxygen release microspheres.** The oxygen release microspheres were fabricated using co-axial electrospaying technique (Fig. 1). The use of co-axial device allows to fabricate microspheres with core-shell structure. The inner stream was PVP/H$_2$O$_2$ complex, and the outer stream was PLGA solution (5 wt% in dichloromethane). The molar ratio of H$_2$O$_2$ and repeating unit of vinylpyrrolidone was 4.5. This ratio allowed the complex to have enough flowability for electrospaying. During the fabrication, the co-axial device was charged at a voltage of +15 kV. The collector aluminum pan was charged at a voltage of −10 kV. The flow rates for the H$_2$O$_2$/PVP complex and PLGA were 0.2 and 1 mL/h, respectively. After fabrication, the microspheres were freeze-dried for 2 h and stored at −20 °C.

![Figure 13. ROS staining of the infarcted hearts harvested 4 weeks after injection. (A–C) CM-H2DCFDA staining for MI (A); Gel (B); and Gel/O$_2$ (C) groups; and (D) ROS content quantified from images. Scale bar = 50 μm. *p < 0.05.](image-url)
Characterization of oxygen release kinetics. Oxygen release from the microspheres was tested in DPBS supplemented with 1 mg/mL of catalase under 1% O\textsubscript{2} condition. To determine oxygen level, an oxygen-sensitive dye luminophore Ru(Ph\textsubscript{2}phen\textsubscript{3})Cl\textsubscript{2} was used. Besides, an oxygen-insensitive fluorophore (rhodamine-B) was utilized as a reference. These two dyes were entrapped in a polydimethylsiloxane (PDMS) membrane. The advantage of using PDMS is that it allows oxygen to interact with dyes due to its high oxygen permeability, while preventing water to penetrate inside owing to its high hydrophobicity. The PDMS membrane was punched into disks with 6 mm in diameter, and placed into a 96-well plate. Two hundred microliters of DPBS supplemented with 1 mg/mL bovine catalase was then added to each well. The plate was placed in a hypoxic incubator (1% O\textsubscript{2}) to balance the

Figure 14. Echocardiographic analysis of Sham, MI, Gel, and Gel/O\textsubscript{2} groups. (A) Ejection fraction; (B) Fractional shortening; (C) Fractional area change. *p < 0.05.
oxygen level in the DPBS. After 24 h, 50 mg of microspheres were quickly added to each well under 1% O₂ environment, and the plate was sealed. The release kinetics was tested in the 1% O₂ incubator at 37 °C (n = 5 for each group). Oxygen release was measured for 4 weeks. At each time point, the fluorescence intensities were measured at 610 nm for Ru(Phen)_2Cl_2 and 576 nm for rhodamine B (excited at 543 nm) by a fluorescent plate reader. The oxygen level was determined according to calibration curve.

**Culture of cardiac fibroblasts, cardiomyocytes and endothelial cells.** Rat cardiac fibroblasts (Cell Applications, San Diego, CA) were cultured using DMEM supplemented with 10% FBS and 1% penicillin-streptomycin. The cells were incubated under 21% O₂, 5% CO₂, and 37 °C conditions. Human umbilical vein endothelial cells (HUVECs) were purchased from Lonza (Walkersville, MD). EGM growth medium supplemented with Endothelial Cell Growth Kit (ATCC, Manassas, VA) was used for culture. The cells between passages 4–7 were used for the studies. Rat neonatal cardiomyocytes (Lonza) were cultured in high-glucose DMEM medium supplemented with 10% FBS and 1% penicillin-streptomycin.

**Characterization of the effect of oxygen release on cardiac fibroblast, cardiomyocyte and endothelial cell survival under hypoxic conditions.** To deliver oxygen release microspheres, they were encapsulated into the above synthesized hydrogel. The hydrogel solution (4 wt%) was mixed with bovine catalase (1 mg/mL) followed by sterilization under UV light for 30 min. The freeze-dried oxygen release microspheres were sterilized separately under UV light. The hydrogel solution and the microspheres (50 mg/mL) were mixed right before injection into the 3D collagen model encapsulated with cells to investigate the effect of oxygen release on cell survival under hypoxic conditions. A 3D collagen model was used to mimic the scenario when the oxygen release system is injected into myocardium after MI. In brief, bovine type I collagen was dissolved in acetic acid solution. The formed collagen solution was then mixed with DMEM supplemented with 10% FBS, and 2 million/mL cells. The mixture was finally neutralized by 0.1 M NaOH. The final collagen concentration was 2 mg/mL. The neutralized mixture was added into 48-well plates (500 µL/well). The plates were placed in a 37 °C incubator for gelation for 45 min to form a construct. The oxygen release system (hydrogel loaded with 50 mg/mL oxygen release microspheres, and 1 mg/mL catalase), or hydrogel alone was then injected into constructs with 50 µL/injection. A total of 5 injections were performed relatively uniformly in each construct. The constructs were cultured in an incubator with 1% O₂. The constructs cultured under normal oxygen condition were used as control.

Cell survival was characterized after 48 h of culture. At least 4 constructs were collected for each experimental group, and digested by papain solution at 60 °C for 24 h. The double-stranded DNA (dsDNA, for live cells) content was measured by PicoGreen assay (Invitrogen). The dsDNA content under hypoxia condition was normalized to that in the normal condition. To image live cells, they were labeled with a live cell tracker CMFDA (5-chloromethylfluorescein diacetate) before seeding into collagen constructs. After 48 h of culture, the cells were imaged using a confocal microscope (Olympus FV1000). At least 3 constructs were imaged for each experimental group.

Cell ROS content was characterized to determine whether oxygen release increased cell oxidative stress that may cause cell death. In brief, cells were trypsinized from the culture flasks after CM-H₂DCFDA (5 µM in PBS) staining for 30 min. The cells were then seeded into the collagen gel using the method described above. After 48 h of culture under 1% O₂ condition or normal condition, samples were harvested, and the cells were imaged by a confocal microscope (Olympus FV1000). Z-stack images were taken in the 30 µm thickness. Four constructs were used for each experimental group. At least 4 different areas were imaged for each construct. All of the images were analyzed by Image J software, the mean gray value in each image was obtained for quantification.

**Characterization of the effect of oxygen release on cardiac fibroblast differentiation into myofibroblast under hypoxic conditions.** To investigate whether oxygen release had impact on cardiac fibroblast differentiation into myofibroblast, the oxygen release system was injected into the 3D collagen model described above. This model mimics the in vivo scenario when the oxygen release system is injected into infarcted heart tissue. In addition, the collagen itself in this model does not change cardiac fibroblast phenotype for at least 48 h. Cardiac fibroblasts were seeded into the collagen gel at the final cell density of 2 million/mL. The constructs were cultured in the culture medium with the addition of TGFβ1 (5 ng/mL) under 1% O₂ condition.

The effect of oxygen release on cardiac fibroblast phenotype was characterized at the mRNA level by real-time RT-PCR, and at the protein level by immunohistochemical analysis. RNA of the samples was isolated using TRIzol. cDNA was synthesized at 1 µg/reaction using a High Capacity cDNA Reverse Transcription kit (ABI). The primer sequence (forward and reverse), melting temperature, and expected product sizes are listed in Table 1. Real-time RT-PCR was conducted in quadruplicate for each sample using SYBR Green master mix on LightCycler® 480 System (Roche). β-Actin was used as the internal reference. Fold of increase was calculated using ΔΔCt method. To identify myofibroblasts formed during culture, immunohistochemical staining of alpha smooth muscle actin (αSMA) was performed for constructs following our established protocol. In brief, the constructs were fixed with 4% paraformaldehyde solution, blocked by 10% goat serum, and permeabilized with 0.1% Triton X-100. The samples were then counterstained with αSMA antibody (mouse-anti-rat) and incubated overnight at 4 °C, followed by staining with secondary antibody Alexa Fluor 546 goat-anti-rat IgG for 1 h. Hoechst 33342 was utilized to stain cell nuclei. The images were taken with a confocal microscope (Olympus FV1000). The αSMA positive myofibroblast density was quantified from at least 5 images for each group.

**Myocardial infarction and implantation of oxygen release system.** All animal experiments were conducted in accordance with the National Institutes of Health Guide for handling laboratory animals and the protocol approved by the Institutional Animal Care and Use Committee of the Ohio State University.
**Characterization of cardiac function by echocardiography.** The rats were subjected to echocardiographic analysis 4 weeks after surgery following our previously established protocol. Baseline was taken prior to the surgery. M-mode echocardiographic and two dimensional images were taken and analyzed using Vevo 2100 High-Resolution *in vivo* imaging system and MS400 transducer (VisualSonics, Toronto, ON, Canada). Images were acquired in a parasternal short and long axis. The dimensions of the left ventricle were measured in short axis view during diastole and systole according to the echocardiographic images.

**Histological analyses.** Four weeks after injection, rat hearts were harvested. The hearts were then underwent retrograde perfusion with DPBS to rinse out the blood followed by fixing in freshly prepared 4% paraformaldehyde overnight. The sections obtained at the level of injection site were chosen for histological studies. The cardiac tissue was embedded in paraffin and sequentially sectioned along the longitudinal axis with 4 μm for each section. Sections were stained with hematoxylin and eosin (H&E), and Picrosirius red (PSR) for wall thickness and collagen composition study, respectively. The thicknesses of at least 10 different areas were measured from each heart sample using ImageJ software. The relative wall thickness was defined as the ratio of myocardium thickness at infarcted area to that of normal area. Type I and III collagens were determined by PSR staining images as yellowish and greenish respectively using polarized light microscopy with appropriate band-pass filters. The percent area of collagen within the remote and MI regions was determined from images (n ≥ 10) using threshold image analysis with ImageJ software.

For immunohistochemical staining, paraffin embedded cardiac samples were sectioned at 4 μm thickness. The sections were counter stained with F4/80, Myosin Heavy Chain (MHC), Ki67, αSMA, and von willebrand factor (vWF) antibodies, respectively. Images (n ≥ 5) of the infarcted area were analyzed. Macrophages were F4/80+ cells. Its percentage was calculated as number of F4/80+ cells over total number of cells in each image. Surviving cardiomyocytes and proliferating cells in the infarcted area were identified as MHC+ and Ki67+ cells, respectively. Myofibroblasts were αSMA+ spindle shaped cells that were not co-localized with vWF+ endothelial cells. Blood vessels were recognized as vWF positive lumen. Densities of MHC+ cells, Ki67+ cells, myofibroblasts, and blood vessels were calculated as #/mm².

To assess ROS content, frozen cardiac tissue was sectioned with 10 μm thickness, and the sections were stained with CM-H2DCFDA. Mean gray value of each cell with positive expression was measured using ImageJ software.

**Statistical analysis.** Statistical analysis was performed using JMP software. One way ANOVA with post-hoc Tukey-Kramer HSD test was utilized for data analysis. Data was presented as mean ± standard deviation. Statistical significance was defined as p < 0.05.

**Data availability.** The datasets generated during and/or analyzed during the current study are available from the corresponding author on reasonable request.

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Author Contributions

Z.F. and J.G. designed the project. Z.F., Z.X., H.N., and N.G. conducted experiments. Y.G., C.L., Y.D., X.C., H.L., X.Z., P.L. and J.M. provided help during experiments. Z.F., X.L.L., Y.D. and J.G. analyzed data. Z.F. and J.G. wrote the manuscript. All authors reviewed the manuscript.

Additional Information

Compeing Interests: The authors declare that they have no competing interests.
