Reactive Oxygen Species (ROS)-Responsive Nanomedicine for Solving Ischemia-Reperfusion Injury

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Ischemia-reperfusion injury (IRI) is a severe condition for most organs, which could occur in various tissues including brain, heart, liver, and kidney, etc. As one of the major hazards, reactive oxygen species (ROS) is excessively generated after IRI, which causes severe damage inside tissues and further induces the following injury via inflammatory response. However, current medical strategies could not thoroughly diagnose and prevent this disease, eventually leading to severe sequelae by missing the best time point for therapy. In the past decade, various nanoparticles that could selectively respond to ROS have been developed and applied in IRI. These advanced nanomedicines have shown efficient performance in detecting and treating a series of IRI (e.g., acute kidney injury, acute liver injury, and ischemic stroke, etc.), which are well-summarized in the current review. In addition, the nano-platforms (e.g., anti-IL-6 antibody, rapamycin, and hydrogen sulfide delivering nanoparticles, etc.) for preventing IRI during organ transplantation have also been included. Moreover, the development and challenges of ROS-responsive nanomedicine are systematically discussed for guiding the future direction.

Keywords: bioresponsive nanomedicine, ischemia-reperfusion injury, reactive oxygen species, organ transplantation, nanoparticles

INTRODUCTION

Instead of restoring and saving tissues from hypoxic conditions, the reperfusion of blood would cause severe damage to those ischemic organs via oxidative stress, which is referred as ischemia-reperfusion injury (IRI). Normally, IRI is strongly associated with a series of diseases, such as the infraction of organs, including stroke, renal infraction, and trauma, etc. Notably, the concern of IRI often affects the best timing of reperfusion, the most effective therapy for treating ischemic disease, for example, the acute myocardial infarction (AMI) (Benjamin et al., 2017). Moreover, the acute myocardial IRI can lead to coronary heart disease (CHD) that accounts for over 366,800 deaths per year and more cases with disability (Benjamin et al., 2018).

In addition to the interruption of cellular homeostasis that could damage cells via apoptotic and autophagic pathways, ischemia-reperfusion would bring great oxidative stress and further injure organs via the generation of redox free radicals, for instance, reactive oxygen species (ROS) (Zhang et al., 2007). Although appropriate ROS is essential for intracellular signaling and immune response against pathogens, the excessive ROS can not be entirely scavenged by cellular antioxidants, which would mediate the severe damage on cells by destroying organelle, cell membrane, and nuclei DNA (Nita and Grzybowski, 2016). To protect injured tissues and prevent further damage of organs during transplantation, the administration of antioxidant reagents for scavenging ROS has been...
IRI and Its Threats

Organs will become ischemic if related vascularst were blockaged, which could induce irreversible damage inside the organs. However, the acute reperfusion for ischecms would cause IRI in some patients, which has been widely observed in various diseases including acute coronary syndrome, hepatic/renal acute injury and organ transplantation (Varadarajan et al., 2004). Eventually, this unexpected complication, IRI will directly lead to a series pathological changes that manifests in brain edema, infarct progression, hemorrhagic transformation, and worsening neurologic system (Zhai et al., 2013; Prabhakaran et al., 2015). More importantly, the severe IRI may further induce systemic diseases including the inflammatory response syndrome (SIRS) and multiple organ dysfunction syndrome (MODS), and increase the mortality of patients, especially for those from the intensive care unit (Cryer, 2000).

General Strategies for Dealing IRI

How to deal with the IRI is of clinical importance, therefore arousing the interests of diverse scientific communities. From the aspect of clinical view, three key factors have been focused on improving the efficiency and safety of reperfusion treatment, including developing different generation thrombectomy devices, lowering the time to reperfusion and applying potential imaging techniques for enrolling suitable patients (Prabhakaran et al., 2015). However, the reality of the synergetic role of various biological mechanisms in IRI made it challenging to prevent or treat. Indeed, some medications with predicted benefits to IRI have not been proved with a definite role in prevention or treatment, likely vitamin C (Hill et al., 2019), morphine (Le Corvoisier et al., 2018), etc. In the field of liver transplantation, nearly all randomized clinical trials have ultimately failed to ameliorate liver IRI in patients (Zhai et al., 2013). Therefore, innovative strategies with smart features for targeting local environmental factor, such as ROS would be an exciting direction for IRI treatment in the future.

Nanomedicine Applied in IRI Therapy

Ischemia could be induced via physical conditions (e.g., elevated blood pressure and diabetes, etc.) or damage, which can further trigger following IRI in various organs, ranging from cerebral to renal system. Notably, the elevated ROS after reperfusion mainly contributes to cellular and tissue damage. Recently, a series of novel nanomedicine (e.g., POM and Ceria NPs, etc.) has been designed for treating affected tissues via the smartly ROS-responsive scavenger and imaging of redox in the ischemic area (Table 1).

Nanomedicine for Cerebral IRI

Cerebral ischemia that generally caused by stroke is associated with high mortality and disability rate. More importantly, ROS produced by cerebral ischemia-reperfusion can further damage brain tissues via neurodegeneration. To overcome this severe disease, various of nanomaterials including polymer NPs (Reddy and Labhasetwar, 2009; Petro et al., 2016; Ghosh et al., 2017; Mei et al., 2019; Mukherjee et al., 2019), ceria NPs (Zhang et al., 2018), polyoxometalate (POM) nanoclusters (Li et al., 2019a), and Framework Nucleic Acids (Li et al., 2019b) have been applied as protective agents, indicating the high efficiency in saving cerebral tissue via ROS scavenging. Notably, the high biocompatibility allows the polymer to be widely recruited for delivering antioxidant reagents (e.g., superoxide dismutase (Reddy and Labhasetwar, 2009), antioxidants catalas (Petro et al., 2016), and curcumin (Mukherjee et al., 2019), etc). Besides, a dual functional polymer NPs, t-PA@iRNP was successfully employed to induce thrombolytic and antioxidant therapies in cerebral tissue (Mei et al., 2019). With the encapsulation of tissue plasminogen activator (t-PA) and conjugation of 4-amino-2,2,6,6-tetramethylpiperidine-1-oxyl (4-amino-TEMOPO), self-assemble t-PA@iRNP (~50 nm) demonstrated an acidic-triggered (pH = 6.2) thrombolytic activity and a significant decrease of ROS production in middle cerebral artery occlusion (MCAO) model mice. More importantly, the antioxidant effect of t-PA@iRNP efficiently avoided the subarachnoid hemorrhage induced by t-PA, releasing a great potential of this dual therapy via synergistic effect.

In addition, novel bio-responsive nanomaterials containing the different status of metal ion (i.e., Mo O and Mo O ) or oxygen-sensitive bio-structure (i.e., DNA) showed excellent capabilities as antioxidants (Li et al., 2019a,b). For instance, a
TABLE 1 | The representatives of novel nanomedicine for treating various IRI via ROS scavenging.

| Application | Type of materials | Nanomedicine | Size (nm) | Mechanism | Investigation | References |
|-------------|------------------|--------------|-----------|-----------|--------------|------------|
| Cerebral IRI | FNA | ac5a-FNA | 10 | Antioxidant (FNA)/C5a-blocking | Protection for cerebral I/R | Li et al., 2019b |
| Cerebral IRI | Polyoxometalate | POM | ~1 | Antioxidant | Protection for cerebral I/R | Li et al., 2019a |
| Cerebral IRI | Polymer | t-PAA@RNP | 48 ± 2 | Antioxidant (LMW nitrooxide)/Thrombolysis | Protection for Middle cerebral artery occlusion | Mei et al., 2019 |
| Cardiac IRI | Polymer | PEG-b-PPS | ~100 | Antioxidant (ginsenoside Rg3) | Protection for Myocardial IRI | Li et al., 2020 |
| Cardiac IRI | Polymer | CLP NPs | 150 ± 18 | Enhanced FL signal via oxidative stress | IVIS imaging of Myocardial IRI | Ziegler et al., 2019 |
| Cardiac IRI | Polymer | MCTD-NPs | 95 | Antioxidant (TPP) | Protection for Myocardial IRI | Cheng et al., 2019 |
| Hepatic IRI | Ceria | Ceria NPs | 4.48 ± 0.8 | Antioxidant | Protection for Partial hepatic IRI | Ni et al., 2019 |
| Hepatic IRI | Ceria | CeO₂ | 10–30 | Antioxidant | Protection for Partial hepatic IRI | Manne et al., 2017 |
| Hepatic IRI | Polymer | BRNPs | 100 | Antioxidant (Bilirubin) | Protection for Partial hepatic IRI | Kim et al., 2017 |
| Hepatic IRI | Polymer | PVO | 550 | Antioxidant (peroxalate esters)/Gas generation | Protection for and US imaging Partial hepatic IRI | Kang et al., 2016 |
| Renal IRI | Carbon QDs | SeCQDs | 40 | Antioxidant (doping selenium) | Protection for Glycerol/Cisplatin-induced AKI Models | Rosenkrans et al., 2020 |
| Renal IRI | FNA | DONs | 90, 120, 400 | Antioxidant | Protection for Glycerol-induced AKI Model | Jiang et al., 2018 |
| Renal IRI | Melanin | MMPP NPs | 4.5 | Antioxidant | Protection for Glycerol-induced AKI Model | Sun et al., 2019 |
| Cardiac IRI | Silica | DATS-MIONs | 230 ± 35 | H₂S delivery (DATS) | Protection for Myocardial IRI | Wang et al., 2019 |
| Organ transplant | Polymer | Anti–IL-6–PLGA-NPs | 100 | Anti–IL-6 delivery | Protection for Heart allografts | Sohjou et al., 2017 |
| Organ transplant | Micelles | TRaM | 15.3 ± 2.3 | Rapamycin delivery | Protection for Aortic interposition allografts | Zhu et al., 2018 |

FNA, Framework nucleic acid; DONs, DNA origami nanostructures; FL, Fluorescence; TPP, Triphenylphosphine; LMW, Low molecular weight; IVIS, in vivo fluorescence imaging system; AKI, Acute kidney injury; PEG, Poly ethylene glycoland; PPS, Poly propylene sulfide.

POM nanocluster was developed and employed as a ROS-responsive agent in treating rats with cerebral I/R injury (Li et al., 2019a). After intrathecal injection, POM could efficiently accumulate in the brain and reduce the oxidative pressure, eventually suppressing the apoptosis of cerebral cells, edema, and infarct volume within the brain (for about 40~50% decrease). Accompanied by IRI, excessive inflammation plays as another key factor for neuronal damage. Recently, a framework nucleic acids (FNA) conjugating with anti-C5a aptamer (ac5a) was applied in the therapy against IRI in the brain (Figure 1A) (Li et al., 2019b). More specifically, the ac5a-FNA could effectively reduce the C5a receptor on polymorphonuclear neutrophils (PMNs) and microglia, eventually bringing down the chemotaxis induced by C5a. Meanwhile, the concentrations of C5a (within plasma and penumbra) and detrimental ROS in cerebral issues were well-scavenged by ac5a-FNA administration, which efficiently protects brain cells from IRI, with a much smaller infarct volume of 17.0 ± 5.6% in comparison with that in control group (41.6 ± 7.1%).

NANOMEDICINE FOR CARDIAC IRI

Among the leading causes of death, cardiovascular disease (CVD), especially the myocardial infarction, is extremely dangerous for any patient. As the biggest challenge, the prevention of IRI greatly affects the outcome of therapy for myocardial infarction. To date, several ROS-responsive nanomaterials have been recruited in cardiac IRI therapy via the delivery of antioxidants (i.e., resveratrol and Ginsenoside Rg3, etc.) (Bae et al., 2016; Cheng et al., 2019; Li et al., 2020). This year, a self-assemble nanoparticle poly ethylene glycol-b-poly propylene sulfide (PEG-b-PPS) that smartly responses to ROS was successfully synthesized and used for ginsenoside Rg3 (Rg3) delivery (Li et al., 2020). PEG-b-PPS-Rg3 exhibited a ROS-dependent release of Rg3 (80% release of Rg3 within 1 day when 1.0 mM ROS was presented) and induced antioxidative stress effects by a strong interaction with FoxO3a (~6.0 kcal/mol). More importantly, the administration of PEG-b-PPS-Rg3 successfully increased the survival rate of I/R rat (90%) in comparison with the group injected with Free Rg3 (40%), indicating its high efficiency in responding to and scavenging cardiac-generated ROS. It is notable that precision targeting may increase therapeutic efficiency. To achieve that, Cheng et al. further modified Szeto-Schiller (SS)-31 and ischemic myocardium targeted peptide (IMTP) for preventing myocardial IRI via mitochondria-targeted antioxidant effect (Cheng et al., 2019). As-prepared MCTD-NPs specifically deliver resveratrol into cardiac tissues, reducing
sizes of the infraction and protecting cardiomyocyte from ROS damage.

Notably, the elevated intracellular Ca\(^{2+}\) is strongly associated with the ROS generation in cardiac cells, which provides another therapeutic strategy via the intervention of Ca\(^{2+}\) influx (via anti-L-Type Ca\(^{2+}\) Peptide and CaMKII inhibitor) (Hardy et al., 2015; Wongrakpanich et al., 2017). However, the efficiency of channel blockade is relatively lower. Thus, the combination of antioxidant (i.e., curcumin) reagents and mitochondria-targeting functionalization (i.e., TPP) are required for improving the therapeutic effect.

Besides, several ROS-responsive nano-platforms were also developed for monitoring the status of IRI during the therapy (Yang et al., 2016; Ziegler et al., 2019). Specifically, Peter Lab developed CLP nanoparticles that could smartly light up the ROS-generated myocardium area after ischemic/reperfusion (Ziegler et al., 2019). The presentation of ROS could restore the fluorescent intensity of Ce6 that was attenuated by the aggregation-induced fluorescence quenching. By this strategy, damaged cardiac areas could be real-time detected and monitored throughout the first day after reperfusion.

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**NANOMEDICINE FOR HEPATIC IRI**

Liver, one of the major organs has played a key role in metabolisms and been involved in various functions, which demands a lot of blood supply. Hepatic ischemia and IRI, a serious complication caused by liver diseases or transplantation would directly trigger irreversible damage in liver tissue. In comparison with other organs, especially the brain, the liver is able to actively “uptake” foreign agents from blood, which allows a series of therapeutic agents to accumulate inside. Thus, antioxidant gene delivery (i.e., expression genes of extracellular superoxide dismutase, catalase or TLR-4 siRNA) (He et al., 2006; Jiang et al., 2011), antioxidant prodrug (i.e., BRAP, etc.) (Lee et al., 2015), and nanoparticles (CeO\(_2\) and platinum NPs, etc.) (Katsumi et al., 2014; Kang et al., 2016; Li et al., 2016; Kim et al., 2017; Manne et al., 2017; Ni et al., 2019) were recruited in treating herpetic IRI. For example, a nanoparticle was designed to release \(\text{p-hydroxybenzyl alcohol (HBA) when } \text{H}_2\text{O}_2 \text{ or LPS was presented, which could remove ROS and decrease the level of pro-inflammation factors (i.e., NO and TNF-\(\alpha\)) (Lee et al., 2015). Moreover, Kang et al. (2016) developed smartly bio-responsive nanoparticles that could provide ultrasound (US) imaging signal...
during H$_2$O$_2$ scavenge. As-prepared PVO nanoparticles could quickly react with ROS and generate CO$_2$ in liver tissue with IRI, offering detectable US signals as early as 25 min post injection.

Additionally, metal nanomaterials exhibiting high reducibility were used as therapeutic agents for hepatic IRI. Among all, CeO$_2$ NPs demonstrated high biocompatibility and efficiency. More specifically, a uniform-sized ceria NPs (∼4.5 nm) was synthesized, containing two valences of Ce (32.01% of Ce$^{3+}$ and 67.99% of Ce$^{4+}$) (Ni et al., 2019). Administration of CeO$_2$ NPs greatly attenuated the IRI in liver tissue as early as 12 h post reperfusion (Figures 1B–D). It is crucial that the ROS scavenge mediated by CeO$_2$ NPs efficiently inhibits the pro-inflammation cytokine secretion (i.e., IL-12, IL-1, and TNF-α, etc.) and an influx of immune cells (i.e., monocyte and neutrophils). Meanwhile, CeO$_2$ NPs also illustrated a high quality in preventing intestinal cells from ROS-based IRI (Gubernatorova et al., 2017).

**NANOMEDICINE FOR RENAL IRI**

Acute kidney injury (AKI) that is triggered by diseases or physical damage could lead to renal disorder/failure. In addition to general therapies like dialysis, ROS-responsive nanomedicine may provide another approach for AKI treatment. Mainly, the functional incapacitation of kidney will directly result in passive accumulation of “waste” inside, including nanomedicine injected. By taking this shortcut, advanced nanomaterials with additional functions such as acid-activated (Yoshitomi et al., 2011) or magnetic resonance imaging (MRI) (Sun et al., 2019) were designed for dealing AKI via ROS reduction. Moreover, Minami et al. also synthesized a nanomedicine (e.g., APP-103) that could effectively accumulate in non-functional kidneys and protect them from IRI during AKI or transplantation (Minami et al., 2020).

In comparison, Cai’s lab successfully developed two nanomedicines specifically targeting to the kidney. Both nanoparticles, SeCQDs and DONs (DNA origami nanostructures) predominantly accumulated in the healthy murine kidney with ∼10 and ∼27%ID/g, respectively, while the kidney after AKI further enhanced the accumulation of nanoparticles (Figure 1E) (Jiang et al., 2018; Rosenkrans et al., 2020). With oxidant-sensitive structures (e.g., doped selenium and nuclei acid), SeCQDs and DONs efficiently removed excessive ROS from renal cells (Figures 1F,G).

**ALTERNATIVE STRATEGIES FOR IRI THERAPY AND ORGAN TRANSPLANTATION**

In addition to an active reduction of cellular ROS, cytoprotective reagents such as hydrogen sulfide (H$_2$S) are also applied as an anti-inflammation agent in various medical situations associated with ischemic diseases. It is notable that various smart nanomedicines have been prepared for H$_2$S delivery, showing promising efficiency (Chen et al., 2019). More specifically, as-designed mesoporous iron oxide and silica nanoparticles could effectively carry and deliver H$_2$S donor diallyl trisulfide (DATS) to ischemic organs (i.e., heart and aortic graft endothelium), eventually reducing the flowing injury caused by reperfusion (Wang et al., 2016, 2019).

Meanwhile, nano-platforms conjugating with immunomodulators could mediate desirable protection for injured or transplanted organs as well. For instance, the modification of antibodies on PLGA NPs successfully moderated the chronic rejection of transplanted heart by targeting IL-6 secreted by allograft-derived dendritic cells (ADDCs) (Solhjou et al., 2017). Furthermore, the endothelial cell inflammation and pro-inflammatory cytokines secretion following tracheal or aortic allografts could be greatly attenuated by the pre-treatment of rapamycin (Nadig et al., 2015; Zhu et al., 2018).

**CONCLUSION/FUTURE DIRECTIONS**

ROS generated by ischemic reperfusion plays a crucial role in the following tissue injury. In other words, this also provides an efficient approach for dealing and preventing IRI via the reduction of ROS. Novel nanoparticles that smartly respond or consume ROS have been designed and proved for their promising performance. However, several directions and challenges should be aware for the future design of ROS-responsive nanomedicine. (a) A combination of multiple functions (i.e., imaging and therapy) in single nanomedicine is highly recommended for monitoring and treating IRI diseases; (b) Desirable IRI therapy may be achieved via dual therapy of ROS scavenge and immune-repress (i.e., H$_2$S delivery); (c) Although alternative strategies, such as antioxidant gene delivery, may provide another direction for IRI therapy, the quick response and antioxidant effects are still the key issues for the future design of ROS-responsive nano-platforms; (d) Tissue-depended design is strongly recommended as well. For example, the antioxidant capability is more important than tissue targeting in AKI model (e.g., passive accumulation in AKI); (e) Sub-organ/cell-targeting for IRI therapy is highly suggested (i.e., antibody-induced cell targeting). Specifically, the most of antioxidant nanoparticles would be internalized by Kupffer cells during the liver IRI, which may lower the therapeutic effect of nanomedicine in major hepatic cells such as liver sinusoidal endothelial cells; (f) The antioxidant efficiency and biocompatibility should be balanced. Thus, advanced ROS-responsive nanomedicine have shown high-performance in dealing with various IRI, which exhibits a great potential in clinical translation, especially organ transplantation.

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WC planned and wrote this review with the help of DL. All authors contributed to the article and approved the submitted version.

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Qiao, Y., He, J., Chen, W., Yu, Y., Li, W., Du, Z., et al. (2020). Light-activatable synergistic therapy of drug-resistant bacteria-infected cutaneous chronic wounds and nonhealing keratitis by cupriferous hollow nanoshells. ACS Nano 14, 3299–3315. doi: 10.1021/acsnano.9b08930

Reddy, M. K., and Labhasetwar, V. (2009). Nanoparticle-mediated delivery of superoxide dismutase to the brain: an effective strategy to reduce ischemia-reperfusion injury. FASEB J. 23, 1384–1395. doi: 10.1096/f.08-116947

Rosenkrans, Z. T., Sun, T., Jiang, D., Chen, W., Barnhart, T. E., Zhang, Z., et al. (2020). Selenium-doped carbon quantum dots act as broad-spectrum antioxidants for acute kidney injury management. Adv. Sci. 7:2000420. doi: 10.1002/advs.202000420

Solhjou, Z., Uehara, M., Bahmani, B., Maarouf, O. H., Ichimura, T., Brooks, C. R., et al. (2017). Novel application of localized nanodelivery of anti-interleukin-6 protects organ transplant from ischemia–reperfusion injuries. Am. J. Transpl. 17, 2326–2337. doi: 10.1111/ajt.14266

Sun, T., Jiang, D., Rosenkrans, Z. T., Ehlerding, E. B., Ni, D., Qi, C., et al. (2019). A melanin-based natural antioxidant defense nanosystem for theranostic application in acute kidney injury. Adv. Funct. Mater. 29:1904833. doi: 10.1002/adfm.201904833

Varadarajan, R., Golden-Mason, L., Young, L., McLaughlin, P., Nolan, N., McEntee, G., et al. (2004). Nitric oxide in early ischaemia reperfusion injury during human orthotopic liver transplantation. Transplantation 78, 250–256. doi: 10.1097/01.TP.0000128188.45553.8C

Wang, W., Liu, H., Lu, Y., Wang, X., Zhang, B., Cong, S., et al. (2019). Controlled-releasing hydrogen sulfide donor based on dual-modal iron oxide nanoparticles protects myocardial tissue from ischemia–reperfusion injury. Int. J. Nanomed. 14:875. doi: 10.2147/IJNN.186225

Wang, W., Sun, X., Zhang, H., Yang, C., Liu, Y., Yang, W., et al. (2016). Controlled release hydrogen sulfide delivery system based on mesoporous silica nanoparticles protects graft endothelium from ischemia–reperfusion injury. Int. J. Nanomed. 11:3255. doi: 10.2147/IJNN.S104604

Wongrakpanich, A., Morris, A. S., Geary, S. M., Mei-Ling, A. J., and Salem, A. K. (2017). Surface-modified particles loaded with CaMKII inhibitor protect cardiac cells against mitochondrial injury. Int. J. Pharm. 520, 273–283. doi: 10.1016/j.ijpharm.2017.01.061

Yang, L., Ren, Y., Pan, W., Yu, Z., Tong, L., Li, N., et al. (2016). Fluorescent nanocomposite for visualizing cross-talk between microRNA-21 and hydrogen peroxide in ischemia-reperfusion injury in live cells and in vivo. Anal. Chem. 88, 11886–11891. doi: 10.1021/acs.analchem.6b03701

Yoshitomi, T., Hirayama, A., and Nagasaki, Y. (2011). The ROS scavenging and renal protective effects of pH-responsive nitroxide radical-containing nanoparticles. Biomaterials 32, 8021–8028. doi: 10.1016/j.biomaterials.2011.07.014

Zhai, Y., Petrovsky, H., Hong, J. C., Busuttil, R. W., and Kupiec-Weglinski, J. W. (2013). Ischaemia-reperfusion injury in liver transplantation—from bench to bedside. Nat. Rev. Gastroenterol. Hepatol. 10, 79–89. doi: 10.1038/nrgastro.2012.225

Zhang, T., Li, C.-Y., Jia, J.-J., Chi, J.-S., Zhou, D., Li, J.-Z., et al. (2018). Combination therapy with LXW7 and ceria nanoparticles protects against acute cerebral ischemia/reperfusion injury in rats. Curr. Med. Sci. 38, 144–152. doi: 10.1007/s11596-018-1858-5

Zhang, W., Wang, M., Xie, H., Zhou, L., Meng, X., Shi, J., et al. (2007). Role of reactive oxygen species in mediating hepatic ischemia-reperfusion injury and its therapeutic applications in liver transplantation. Transpl. Proc. 39, 1332–1337. doi: 10.1016/j.transproceed.2006.11.021

Zhu, P., Atkinson, C., Dixit, S., Cheng, Q., Tran, D., Patel, K., et al. (2018). Organ preservation with targeted rapamycin nanoparticles: a pre-treatment strategy preventing chronic rejection in vivo. RSC Adv. 8, 25909–25919. doi: 10.1039/C8RA01555D

Ziegler, M., Xu, X., Yap, M. L., Hu, H., Zhang, J., and Peter, K. (2019). A self-assembled fluorescent nanoprobe for imaging and therapy of cardiac ischemia/reperfusion injury. Adv. Ther. 2:1800133. doi: 10.1002/adtp.201800133

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