3, 4-dihydroxyl-phenyl lactic acid restores NADH dehydrogenase 1 α subunit 10 to ameliorate cardiac reperfusion injury

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The present study aimed to detect the role of 3, 4-dihydroxyl-phenyl lactic acid (DLA) during ischemia/reperfusion (I/R) induced myocardial injury with emphasis on the underlying mechanism of DLA antioxidant. Male Spragu-Dawley (SD) rats were subjected to left descending artery occlusion followed by reperfusion. Treatment with DLA ameliorated myocardial structure and function disorder, blunted the impairment of Complex I activity and mitochondrial function after I/R. The results of 2-D fluorescence difference gel electrophoresis revealed that DLA prevented the decrease in NDUFA10 expression, one of the subunits of Complex I. To find the target of DLA, the binding affinity of Sirtuin 1 (SIRT1) to DLA and DLA derivatives with replaced two phenolic hydroxyls was detected using surface plasmon resonance and bilayer interferometry. The results showed that DLA could activate SIRT1 after I/R probably by binding to this protein, depending on phenolic hydroxyl. Moreover, the importance of SIRT1 to DLA effectiveness was confirmed through siRNA transfection in vitro. These results demonstrated that DLA was able to prevent I/R induced decrease in NDUFA10 expression, improve Complex I activity and mitochondrial function, eventually attenuate cardiac structure and function injury after I/R, which was possibly related to its ability of binding to and activating SIRT1.

Coronary heart disease is a leading cause of death all over the world, and its incidence is increasing at an alarming rate1. After an acute myocardial infarction, percutaneous coronary intervention has become the most common strategy to restore myocardial perfusion, however it does not reduce the risk of mortality2, due to numerous complications of I/R injury. I/R causes cardiomyocyte death, myocardial stunning and left ventricular remodeling, which contribute to the reduction of cardiac output and myocardial fibrosis leading to the development of heart failure3,4.

One of the major consequences of I/R is mitochondrial dysfunction, resulting in mitochondria-derived myocardium damage by reactive oxygen species (ROS) production, adenosine triphosphate (ATP) reduction, increased mitochondrial permeability, cytochrome c release and activation of programmed cell...

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that CP ameliorates I/R-induced myocardial damage and fibrosis in rats. DLA itself is well known to have cardiovascular diseases by the US Food and Drug Administration in 2013. Our previous results indicated that DLA is safe for use in humans. Therefore, we selected rats with a similar tension and placement of the ligature among the groups (Fig. 1C). The dose of 5 mg/kg was selected as an optimal dose of DLA and applied in all subsequent experiments.

Results

DLA administration diminishes infarct size, preserves myocardium structure and reduces leukocyte infiltration after myocardial I/R. We first examined whether DLA exhibited a cardioprotective role during I/R in rats. Upon I/R, the hearts from DLA treated rats showed a reduced ratio of IA/AAR in a dose-dependent manner compared to those from saline treated rats (Fig. 1A,B). However, the ratios of AAR/LA were the same in all I/R groups regardless of DLA treatment or not, indicating a similar tension and placement of the ligature among the groups (Fig. 1C). The dose of 5 mg/kg was selected as an optimal dose of DLA and applied in all subsequent experiments.

Micrographs of hematoxylin/eosin (H&E) staining sections shown in Fig. 2A revealed that myocardial tissues from rats undergoing I/R were morphologically altered, including disordered arrangement and disruption of myocardial fibers, myocardial interstitial edema and inflammatory cell infiltration. Noticeably, administration of DLA markedly protected hearts from morphological alterations after I/R. F-actin, as a component of thin filament, has an important role in contraction force generation. Confocal microscopy studies showed a pronounced F-actin disarrangement and disruption in myocardium sections after challenge by I/R which were attenuated by DLA treatment (Fig. 2B).

As a marker enzyme of neutrophils, MPO expression in myocardium was assessed by immunohistochemistry. Few cells exhibited MPO-positive staining in SHAM group. On the contrary, I/R evoked a considerable increase in infiltration of neutrophils. Interestingly, the I/R-induced increase in the number of MPO-positive cells in heart was diminished by treatment with DLA (Fig. 2C). The quantitative result of the MPO-positive cells in surrounding infarction areas in myocardium is consistent with the qualitative survey, as shown in Fig. 2D.

DLA administration leads to better preservation of LV function and MBF after I/R. Given to the fundamental role of F-actin in generating contractile force, we then tested whether DLA treatment would benefit cardiac function during I/R. LV function at baseline was comparable among four groups. LV developed pressure (max) fell substantially at 30 min of ischemia compared to sham operation. However, DLA treatment improved LV developed pressure (max) considerably compared with
saline treated group during reperfusion (Fig. 3A). Similar results were observed for LV systolic pressure (Supplementary Fig. S1A). Furthermore, reduction in mean artery pressure (Fig. 3C) and artery systolic pressure (Supplementary Fig. S1B) induced by I/R was notably restored by DLA administration after 90 min of reperfusion. In addition, LV developed pressure (min) rose dramatically at 30 min of ischemia compared with SHAM group, DLA treated rats exhibited better recovery of LV developed pressure (min) than normal saline treated group during reperfusion (Fig. 3B). Consistent with the results of LV developed pressure (min), increase in LV end diastolic pressure (Fig. 3D) caused by I/R was substantially prevented by DLA administration at the end of reperfusion. Taken together, these results demonstrated that the impaired LV function in response to I/R was significantly protected in DLA treated group. The protective role of DLA for heart structure and function was further confirmed by echocardiography analysis conducted at the third week after I/R. As shown in Supplementary Table S3 and Fig. S4, compared with SHAM group, I/R led to an obvious increase in LVIDd, LVIDs, LV Vold and LV Vols, but a decrease in EF%, FS% and FAC%, most of which, except LVIDd and LV Vold, were attenuated by DLA treatment.

Better recovery of left ventricular function suggested that DLA treatment might improve coronary perfusion after I/R. To verify this speculation we then examined MBF by using a Laser Doppler Perfusion Imager. Representative Laser Doppler pictures of MBF at different time points of I/R were acquired from four groups, as illustrated in Fig. 3E. The time courses of quantitative results showed that MBF fell evidently after 30 min of ischemia compared to SHAM+NS groups, however, DLA treatment led to a better MBF recovery in comparison with I/R+NS group (Fig. 3F).

**DLA administration represses I/R-induced apoptosis.** TUNEL staining was performed to investigate the extent of apoptosis in the AAR region in four groups. Confocal images and statistical results of TUNEL staining indicated that the hearts from DLA treated rats showed a significant decrease in the number of TUNEL positive cells compared to that from I/R+NS group after I/R, whereas only few TUNEL positive cells were detected in the hearts from SHAM group (Supplementary Fig. S2A and B). Apoptosis is a process that is controlled by apoptosis related proteins, thus we assessed Bcl-2, Bax and Cleaved-caspase 3 in whole-heart lysates. The results demonstrated that DLA exerted anti-apoptosis

**Figure 1.** DLA administration diminishes I/R-induced infarct. **A**, Representative images of myocardial tissue sections stained with TTC and Evans blue. The rats in I/R groups were subjected to 30 min ischemia and 90 min reperfusion with intravenous injection of normal saline (NS) or DLA (dose ranging from 1.25 mg/kg to 20 mg/kg). The non-ischemic area is indicated by blue, the AAR by red and the IA by white. **B** and **C**, Quantitative analysis of infarct size of myocardial tissues. The ratios of IA to AAR and AAR to LA are shown. Results are presented as mean ± S.E.M (n = 6) * p < 0.05 vs. SHAM + NS group, # p < 0.05 vs. I/R + NS group, † p < 0.05 vs I/R + DLA 1.25 mg/kg, & p < 0.05 vs I/R + DLA 2.5 mg/kg.
role by upregulating the expression of anti-apoptotic protein Bcl-2 and downregulating the expression of pro-apoptotic protein Bax, as compared to rats in I/R + NS group (Supplementary Fig. S2C and D).

Furthermore, ratio of Bcl-2/Bax, a determinant of the susceptibility of myocardium to apoptosis, was noticeably diminished after I/R, which was restored in the presence of DLA (Supplementary Fig. S1C). Consistent with these results, DLA prevented hearts from I/R induced increase in caspase-3 cleavage (Supplementary Fig. S2E).

**DLA administration restores I/R induced reduction in NDUFA10 expression.** 2-D fluorescence difference gel electrophoresis (2D-DIGE) was performed in order to reveal the mechanism by which DLA protects hearts from I/R injury. Representative 2D gel images of all groups are shown in Fig. 4A. Four protein spots were identified as differentially expressed between I/R + NS group and I/R + DLA group (Supplementary Table S1). Among the identified proteins, NDUFA10 was the only protein located in mitochondria. Mitochondria is the major contributor to the oxidative damage and myocardium apoptosis under the conditions of I/R. Considering the role of DLA in antioxidation and anti-apoptosis, we focused on the study of mitochondrial protein NDUFA10.

Selected regions of the 2D gels displaying NDUFA10 protein spot and its three-dimension view in Fig. 4A and B showed that I/R induced a dramatic reduction in NDUFA10 expression in comparison with sham operation, and this reduction was protected by treatment of DLA. Western blot confirmed the results of 2D-DIGE (Fig. 4C). NDUFA10 mRNA levels were also assessed by reverse transcription real-time PCR. Likewise, DLA administration dramatically restored the reduction in NDUFA10 mRNA expression induced by I/R (Fig. 4D).

**DLA administration prevents against the impairment in Complex I function induced by I/R.** Given to the fact that NDUFA10 is essential for proper function of Complex I, we speculated that NDUFA10 decrement would result in dysfunction of Complex I and mitochondria in the heart. To test this hypothesis, mitochondrial Complex I activity was first determined, and the results are shown in Fig. 5A in which OD at 450 nm increased over time with the slope being proportional to Complex I activity.
activity. Evidently, impaired Complex I activity induced by I/R was remarkably reversed by injection of DLA (Fig. 5B).

Increased Complex I activity was expected to improve mitochondrial respiration, resulting in enhanced ATP production and less free radical accumulation. Therefore, energy metabolic conditions and redox state of myocardial tissue were next assessed. Consistent with the preserved Complex I activity, the reduction in myocardial ratio of ATP/ADP and ATP/AMP after I/R was considerably restored by DLA administration (Fig. 5C,D). In addition, I/R induced an increase in the level of malondialdehyde (MDA), an indicator of cellular lipid peroxidation, which was dramatically reduced by administration of DLA (Fig. 5E). A decrease in Complex I activity would impair the mitochondrion themselves. Mitochondrial ultrastructure observed by transmission electron microscopy (Fig. 5F) showed that I/R resulted in distinctive ultrastructure alterations, including disordered mitochondrion distribution with disarranged and obscure crista and vacuoles within the matrix, accompanying by disrupted myofilament and sarcomere. Noticeably, these deleterious effects of IR were markedly attenuated by DLA administration (Fig. 5F).
DLA binds to and activates SIRT1 to preserve NDUFA10 expression and Complex I activity. The question arose was that what was the target upstream NDUFA10 for DLA acting at? To this end, SIRT1 has been reported to induce several mitochondrial respiration chain gene expression, it is thus likely that DLA elevates NDUFA10 mRNA levels through activating SIRT1. To test this hypothesis, BLI was conducted to determine the binding affinity of DLA to SIRT1. As shown in Fig. 6A, DLA was able to bind to SIRT1 in a dose-dependent manner. \( K_D \) derived from recorded BLI curve were \( 3.37 \times 10^{-4} \) (M). To verify the results of BLI, the binding capacity of DLA to SIRT1 was analyzed by SPR assay. Consistent with BLI, results of SPR indicated that DLA bound to SIRT1 in a dose-dependent way (Fig. 6B). \( K_D \) constant of DLA binding to SIRT1 was \( 7.84 \times 10^{-4} \) (M).

DLA having a capacity of directly interacting with SIRT1 led us to ask: what is the impact of DLA on SIRT1 content and activity? As shown in Fig. 6C, DLA protected hearts from I/R-caused reduction in NDUFA10 expression. A, Representative 2D-DIGE images of heart lysates from ischemic region in various groups. Selected and enlarged regions of 2D-DIGE images displaying NDUFA10 are shown in the middle panel, while three-dimensional views of the selected regions displaying NDUFA10 expression alterations shown in the lower panel. Arrows indicate NDUFA10. B, The overlay images with yellow spots indicating proteins that have equal expression levels in both groups, green spots indicating proteins down-regulated in I/R+NS group. Arrows indicate NDUFA10. C, Western blot and statistical analysis of NDUFA10 based on the data of 3 independent experiments normalized to GAPDH. The gels have been run under the same experimental conditions. D, NDUFA10 mRNA levels determined by reverse transcription real-time PCR. Quantitative results of mRNA expression were derived from 5 independent experiments and normalized to GAPDH. Results are presented as mean ± S.E.M. * \( p < 0.05 \) vs. SHAM+NS group, # \( p < 0.05 \) vs. I/R+NS group.
in SIRT1 protein expression. SIRT1 inhibitors sirtinol and EX-527 were applied to determine the effects of DLA on deacetylase activity of SIRT1. Doses of inhibitors were calculated based on the concentration used in vitro experiments. Optimal inhibitor dose was selected from three doses administrated, the dose of sirtinol and EX-527 applied in subsequent experiments was 4.8 mg/kg and 4 mg/kg, respectively. Based on the results of in vivo deacetylase activity detection, impaired deacetylase activity induced by I/R was remarkably reversed by injection of DLA, while preserved deacetylase activity caused by DLA administration during I/R was abolished by SIRT1 inhibitor sirtinol and EX-527 (Fig. 6D). DLA treatment dramatically brought down increased acetylated-Foxo-1 (Ac-Foxo-1) levels after I/R (Supplementary Fig. S3A), which verified the results of SIRT1 activity. Moreover, manganese superoxide dismutase (MnSOD), which could be upregulated by SIRT1 in the transcriptional level29, was obviously decreased after I/R, which was reversed by DLA administration as well (Supplementary Fig. S3B).

Furthermore, the beneficial effects of DLA on restoring NDUFA10 expression (Fig. 6E) and Complex I activity (Fig. 6F) were abolished by sirtinol and EX-527. The result of present study implied NDUFA10 as a downstream effector of SIRT1 activation by DLA administration.

Figure 5. DLA administration restores Complex I activity and mitochondrial function after I/R. A, Time course of OD at 450 nm, the slope being proportional to Complex I activity. B, Complex I activity based on the slope of increase in OD at 450 nm. C and D, Quantitative results of ratios of ATP/ADP and ATP/AMP in ischemic heart homogenates in various groups after I/R, respectively. E, Statistical analyses of MDA of ischemic heart homogenates in various groups after I/R. F, Representative electron micrographs of myocardial tissues of rats from 4 groups. a: mitochondrial swelling. b: disarranged and obscure crista. c: vacuoles among crista. d: disrupted myofilament and sarcomere. Results are presented as mean ± S.E.M (n = 6) * p < 0.05 vs. SHAM+NS group, # p < 0.05 vs. I/R+NS group.
Figure 6. DLA binds to and activates SIRT1. A, Sensorgram binding curves of four different concentrations of DLA (2500, 5000, 10000, and 20000 μM, curves from bottom to top) measured by BLI. B, Representative sensorgrams obtained from the injections of DLA at concentrations of 312.5, 625, 1250, 2500, and 5000 μM (curves from bottom to top) using SPR. C, Representative western blot bands of SIRT1 in various groups shown above and quantitative results below. The gels have been run under the same experimental conditions. D, The deacetylase activity of rats myocardial tissue (n = 6) in various groups with or without SIRT1 inhibitor sirtinol and EX-527. E, The representative western blot bands and the quantitative results of NDUFA10 in various groups with or without sirtinol and EX-527. The gels have been run under the same experimental conditions. Quantification results for NDUFA10 band intensities were normalized to GAPDH from 3 independent experiments. F, Complex I activity based on the slope of increase in OD at 450 nm in all groups with or without sirtinol and EX-527. Results are presented as mean ± S.E.M. * p < 0.05 vs. SHAM+NS group, # p < 0.05 vs. I/R+NS group, & p < 0.05 vs. I/R+DLA group.
**DLA binds to SIRT1 in a phenolic hydroxyl dependent manner.** Then we tried to elucidate the mechanisms of how DLA could bind to SIRT1. DLA derivatives were synthesized by replacing both phenolic hydroxyl groups of DLA into methoxyl (Compound A) or hydroxyl (Compound B) (Fig. 7A). The association constant ($k_a$), dissociation constant ($k_d$), and $K_D$ of DLA, Compound A and B binding to SIRT1 were summarized in Supplementary Table S2, showing that the $K_D$ constants of Compound A and B binding to SIRT1 were 0.638 (M) and 0.123 (M), respectively. $K_D$ values and binding curves of three molecules to SIRT1 shown in Fig. 7B indicated that both Compound A and B exhibited lower affinity than DLA for SIRT1. Furthermore, the beneficial effects of DLA on preventing myocardial infarct during I/R was abolished by replacement of phenolic hydroxyl groups, Compound A and B exhibiting an impaired ability to decrease IR-caused myocardial infarct compared with DLA (Fig. 7C–E). Taken together, these results demonstrated that cardioprotective effect of DLA during I/R was correlated with its binding to and activating SIRT1, a process depending on the presence of phenolic hydroxyl groups.

**SIRT1 is required for the protective role of DLA in increasing NDUFA10 expression and Complex I activity, reducing mitochondrial ROS and cell death.** In order to confirm the involvement of SIRT1 in DLA alleviating I/R-induced cardiomyocyte injury, SIRT1 expression was knocked down by treatment with siRNA in H9c2 cells exposing to hypoxia/reoxygenation (H/R). As shown in Fig. 8A,B, SIRT1siRNA considerably reduced the protein level of SIRT1 under normal or H/R condition. As an important subunit of Complex I, NDUFA10 in H9c2 cells was examined by western blotting assay, as shown in Fig. 8C. As expected, H/R led to a decrease in NDUFA10 while treatment with DLA restrained NDUFA10 expression after H/R. SIRT1 knocked down by siRNA decreased NDUFA10 after DLA treatment in H/R condition. Mitochondrial Complex I activity in H9c2 cells was then determined, and the results are shown in Fig. 8D in which OD at 450 nm increased over time being proportional to Complex I activity. Evidently, increased Complex I activity induced by DLA treatment after H/R was remarkably reversed by SIRT1siRNA. Complex I activity is necessary for mitochondrial respiration, which maintains intracellular normal free radical level. Therefore, we evaluated the basal and inducible mitochondrial ROS level in H9c2 cells using molecular probe DHR 123. As shown in Fig. 8E, the ROS level was considerably increased after H/R compared to control cells. DLA treatment reduced the H/R-elevated ROS level, while SIRT1 knockdown reversed the antioxidative role of DLA, leading to a higher ROS in H9c2 cells. The results of CCK-8 assay (Fig. 8F) showed that at the end of reoxygenation, the viability of H9c2 cells was decreased to approximate a half of control group. Treatment with DLA attenuated the H/R-induced depression in cell survival. DLA treatment of cells with knocked down SIRT1 showed a remarkably reduced cell viability compared with DLA group without SIRT1siRNA. Above results indicated that SIRT1 played an important role in DLA restoring NDUFA10 expression and Complex I activity after H/R, as well as in maintaining intracellular redox state and cell viability.

**Discussion**

Our results demonstrated for the first time that DLA could restore NDUFA10 expression and Complex I activity after I/R, which is probably attributable to the ability of DLA to bind to SIRT1 and preserve SIRT1 activity, which eventually results in reduced infarct size, improved LV function and better recovery of MBF (Supplementary Fig. S5).

Our 2D-DIGE result provided novel insights into the role of mitochondria in DLA mediated anti-apoptotic effect in response to I/R, showing that NDUFA10, among others, was better maintained by DLA after I/R injury. More importantly, restoration of NDUFA10 by DLA was accompanied by an improved mitochondrial Complex I activity after I/R. Under the conditions of hypoxia/reoxygenation30, ischemic and I/R31, a decreased mitochondrial Complex I activity was observed. Given to the central role of Complex I in electron transport and oxidative phosphorylation, restoration of Complex I activity is expected to be one of the most promising strategies for reducing ROS and increasing ATP production after I/R. The results of present study, as shown by the decrease in MDA level and increase in ATP content by DLA, supported the above speculation, suggesting a beneficial role of DLA-provoked Complex I activity.

An emerging concept is that blockade of ETC, including Complex I, during ischemia is cardioprotective32. One possibility accounting for this discrepancy may be that mitochondrial damage caused by ETC mainly occurs in ischemia period, the continuation of enhanced mitochondrial aerobic respiration without oxygen results in deletion of cardiolipin, leading to loss of cytochrome c and the onset of mitochondrial permeability transition, which taken together promote myocardial cell apoptosis32,33. This is why most, if not all, the cardioprotective roles for blockade of Complex I are achieved by intervention immediately before ischemia or during ischemia. In the present study, DLA was administrated 1 min before reperfusion until the end of reperfusion. During this phase, sufficient oxygen is available that requires a dramatic increase in Complex I activity to drive the electron passing through ETC. Another possibility is that DLA imposed effect not directly on Complex I but on SIRT1, which modulated a range of proteins, including NDUFA10. The cardioprotective role of DLA observed in current study was a collective outcome of the activation of these proteins.

Study has demonstrated that SIRT1 mutant mice exhibit notable developmental defects in the heart34. Moderate increased SIRT1 was found to protect the heart against oxidative stress by upregulating...
Figure 7. DLA derivatives exhibit lower affinity for SIRT1 and impaired ability for reducing infarct size after I/R. A, The structures of DLA, Compound A and Compound B. B, SPR binding curves of DLA, Compound A and Compound B to SIRT1, subtracting non-specific binding to bovine serum albumin (BSA), at the same concentration of 1250 μM. C, Representative images of myocardial tissues stained with TTC and Evans blue in rats of SHAM group and rats subjected to I/R with or without injection of DLA (5 mg/kg), Compound A (5 mg/kg) and Compound B (5 mg/kg). D to E, Quantitative analyses of infarct size in myocardial tissues. The ratios of IA to AAR (D), and AAR to LA (E) are shown. Results are presented as mean ± S.E.M (n = 6) * p < 0.05 vs. sham group, # p < 0.05 vs. I/R group, & p < 0.05 vs. I/R+DLA group.
antioxidant expressions. Overexpression of SIRT1 was revealed to prevent myocardium apoptosis in response to serum starvation. Furthermore, cardiac-specific overexpression of SIRT1 was also shown to reduce infarct size and improve left ventricular functions after I/R, whereas cardiac-specific SIRT1 knockout mice were more susceptible to I/R injury. Thus, SIRT1 is considered to be an effective target in alleviating I/R induced injury.

SIRT1 has been reported to increase transcription of antioxidants. Consistent with this finding, we observed that MnSOD content was enhanced in response to the administration of DLA. DLA is well known for its antioxidant activity, however, the underlying mechanisms are poorly understood. The results of present study implied that antioxidative effect of DLA may be attributable to maintenance of Complex I integrity and activity, which prevented electron leakage from ETC, as well as to upregulation of MnSOD that was mediated by DLA binding to and activating SIRT1.

Our results revealed DLA was able to bind to SIRT1 in vitro in a dose dependent manner, possibly depending on the presence of phenolic hydroxyl groups. However, we cannot conclude that phenolic hydroxyl is the exact binding group of DLA to SIRT1. More experiments need to be carried out to investigate the roles of other groups in addition to phenolic hydroxyl. Besides, replacing phenolic hydroxyl of DLA into methoxyl or hydroxyl might lead to an alteration in conformation and physic-chemical property of DLA, which in turn may affect its binding affinity for SIRT1. Especially, structural evidence is required to confirm the binding of DLA to SIRT1, and the binding affinity of DLA to SIRT1 needs to be assessed in vivo. Thus, the exact mechanism responsible for the interaction between DLA and SIRT1 remains to be clarified.

We have identified DLA as a regulator of mitochondrial function probably through modulating SIRT1 activity during I/R injury in the heart. However it remains unclear how SIRT1 achieved the transcriptional regulation of NDUFA10 mRNA after DLA administration. Several lines of evidence suggest that PGC-1α serves as the downstream effector of SIRT1 in upregulating mitochondrial respiration genes.
Thus, PGC-1α might be responsible for upregulating NDUFA10 expression in the presence of DLA treatment. Nevertheless, more works are required to clarify the signaling pathway which links up SIRT1 to NDUFA10 for DLA to attenuate the Complex I dysfunction after I/R.

**Summary.** In the present study, we confirmed the cardioprotective role of DLA against I/R injury and revealed the ability of DLA to bind to and activate SIRT1, resulting in enhanced expression of NDUFA10, Complex I activity and mitochondrial function after I/R. These results provide clue for identification of the signaling pathway thereby DLA exerts its beneficial effects, as well as for designing novel medicine to cope with I/R-induced cardiac injury.

**Methods**

**Animals.** Male Sprague-Dawley (SD) rats weighing 240-260 g were purchased from the Animal Center of Peking University Health Science Center (Certificate no. SCXK (jing) 2006-0008). All experimental procedures involving animals were approved by Peking University Biomedical Ethics Committee Experimental Animal Ethics Branch (LA2010-001), complying with the Guide for the Care and Use of Laboratory Animals (NIH publication no. 85-23, 1996).

**I/R model establishing and drug administration.** Rats were anesthetized by intraperitoneal injection of sodium pentobarbital (60 mg/kg) and maintained by 1% isoflurane inhalation. Anesthetic monitoring was achieved by a pinch on the toe. I/R animal model was established as previously described. In SHAM group, the rats underwent the same procedure, except for tightening the ligature. A polyethylene catheter was inserted into left femoral vein for administration of DLA (Kun Ming Feng-Shan-Jian Medical Company, Yunnan, China), Compound A, Compound B or normal saline (NS). DLA, Compound A and Compound B were dissolved in 2 ml NS, and 0.5 ml was given intravenously 1 min before reperfusion, and the remaining was continuously administrated till the end of reperfusion at a speed of 1 ml/h. DLA was given at a dose of 1.25 mg/kg, 2.5 mg/kg, 5 mg/kg, 10 mg/kg, or 20 mg/kg. Compound A and Compound B were injected at the dose of 5 mg/kg.

**2,3,5-triphenyltetrazolium chloride and Evans blue staining.** At the end of reperfusion, the left descending artery (LAD) was re-ligated, and 2 ml of 4% Evans blue (Sigma Aldrich, Missouri, USA) was injected into the right femoral vein via intravenous catheter to delineate the area at risk (AAR). Then the heart was excised at a 5 mm level above the apex cordis and sliced transversely into five sections (1 mm thick). The slices were incubated in 2% 2,3,5-triphenyltetrazolium chloride (TTC) (AMRESCO, Ohio, USA) at 37 °C for 15 min to identify the infarction area (IA). AAR, IA and left ventricular area (LA) were analyzed by Image-Pro Plus 5.0 software (Media Cybernetic, Maryland, USA).

**Echocardiographic analysis.** The left ventricular wall thickness and heart function were evaluated at the third week after myocardial I/R, using a Vevo 770 High-Resolution Imaging System (Vevo 770, Visual Sonics Inc, Toronto, Canada) with a 17.5 MHz linear array transducer (model 716). The parameters assessed were as follows: left ventricular internal diameter at end-diastole/end-systole (LVIDd/LVIDs), left ventricular anterior wall at end-diastole/end-systole (LVVd/LVVs), left ventricular volume at end-diastole/end-systole (LV Vold/LV Vols), left ventricular mass (LV Mass), left ventricular posterior wall at end-diastole/end-systole (LVPWd/LVPWs), left ventricle ejection fraction (EF%), left ventricle fractional shortening (FS%) and left ventricular fractional area change (FAC%).

**SIRT1 inhibitor administration.** SIRT1 inhibitors sirtinol and EX-527 were injected intravenously 45 min before DLA administration. The dose of sirtinol used in the experiments was 1.2 mg/kg, 2.4 mg/kg, or 4.8 mg/kg, and the dose of EX-527 was 1.2 mg/kg, 4 mg/kg, or 12 mg/kg. SIRT1 inhibitors were first dissolved in stock solutions of DMSO, and for treatment, the final concentration of DMSO was 0.2% (v/v). In groups without inhibitor administration, DMSO at the same final concentration was given to rats. No obvious adverse effects were observed in DMSO treated groups.

**Detection of deacetylase activity.** After ischemic heart tissue was collected and homogenized, proteins were extracted. Tissue Sirtuin activity was detected using a SIRT1 fluorometric assay kit (Enzo, New York, USA), according to manufacturer instruction. Samples with 100 g of total protein were incubated with Fluor de Lys-SIRT1 substrate (100 μM), NAD⁺ (50 μM) at 37 °C for 60 min, and 50 μl mixture of developer and nicotinamide (2 mM) was added by pipetting. Fluorescence kinetics was monitored for 1 h with measuring at 360 nm (excitation) and 460 nm (emission) at 37 °C every 5 minutes.

**Electron microscopy.** After reperfusion, hearts were fixed by infusion with a fixative containing 5% paraformaldehyde and 2% glutaraldehyde for 40 min. Cardiac tissue was excised as described before, and trimmed into 1 mm³ and fixed overnight in 3% glutaraldehyde at 4 °C. Trimmed tissues were post-fixed by 1% osmium tetraoxide for 2 h and prepared for thin sectioning. The thin sections were observed with a transmission electron microscope (JEM 1230, JEOL, Tokyo, Japan).
2-dimension fluorescence difference gel electrophoresis. The samples were harvested as described before and prepared with a 2-dimension (2D)-Clean up Kit (GE Healthcare, Buckinghamshire, UK). Equal amounts of proteins from all samples were pooled together as the internal standard labeled with Cy2. Fifty micrograms of lysates were labeled with 400 pmol of Cy3 or Cy5. The DIGE experimental design was as follows: Gel 01, internal standard (Cy2), SHAM+NS group (Cy3), I/R+NS group (Cy5); Gel 02, internal standard (Cy2), I/R+DLA group (Cy3), SHAM+DLA group (Cy5). Immobilized pH gradient (IPG) strips (24 cm, pH 3-10, and nonlinear) were prepared with labeled samples as previously reported. Then IPG strips were placed on the top of 12% polyacrylamide gels, and electrophoresed for ~10 hours at 30 mA per gel using an Etan DALT Six system (GE Healthcare, Buckinghamshire, UK). Then, protein spots were selected and identified as described previously.

Surface plasmon resonance. Carboxymethylated 5 (CM5) sensor chip was docked into the BLAcore 3000 (GE Healthcare, Buckinghamshire, UK), and prepared as previously reported. Human recombinant SIRT1 (Sigma, Missouri, USA) was immobilized on CM5 sensor chip by injecting 40 μl of SIRT1 (40 μg/ml in 10 mM sodium acetate, pH 4.0) at the rate of 5 μl/min. DLA, Compound A, and Compound B were prepared as a 20 mM solution in running buffer before the experiment, and two fold diluted by running buffer into 10 mM, 5 mM, 2.5 mM, 1.25 mM, 625 μM, 312.5 μM before injection. Analytes were injected at 30 μl/min over SIRT1 and control sensor chip. Equilibrium dissociation (K_D) was calculated by fitting a 1:1 Langmuir model using the BIA evaluation 4.1 software (GE Healthcare, Buckinghamshire, UK).

Biolayer interferometry. All Biolayer interferometry (BLI) experiments were performed on an Octet Red instrument (ForteBio Inc., California, USA) at 25°C. Super Streptavidin (SSA) biosensors were loaded with biotinylated SIRT1 (20 μg/ml) antibody followed by recombinant human SIRT1 (50 μg/ml, 200 μl). Association of DLA (20 mM, 10 mM, 5 mM, 2.5 mM) and SIRT1 was observed by immersing Super Streptavidin biosensors into PBS for 240 s. K_D was analyzed by 1:1 binding global fit model using the Octet data analysis software 6.3 (ForteBio Inc., California, USA).

Cell culture and H/R model. H9c2, a rat cardiac myoblast cell line (ATCC, Maryland, USA), was cultured in Dulbecco’s Modified Eagle’s Medium (Invitrogen, Grand Island, UK) containing 4 mM L-glutamine, 4.5 g/L glucose, and 10% fetal bovine serum (Invitrogen, Grand Island, UK) at 37°C in a humidified incubator with 95% air and 5% CO_2_. The cells were subjected to hypoxia/reoxygenation (H/R) by incubation in a microaerophilic system (Thermo Scientific, Waltham, MA) at 5% CO_2 and 1% O_2 for 12 h followed by normoxic conditions for 6 h with or without 0.1 mM DLA treatment.

Small interfering RNA transfection. When 50%-60% confluence was reached, H9c2 cells were transfected with 20 μM SIRT1 small interfering RNA (siRNA) or nontargeting control siRNA complexed with Lipofectamine 2000 (Invitrogen, Carlsbad, CA), according to manufacturer’s instructions. The sense sequences of SIRT1 siRNA was as follows: 5′- CAGGAAUCCAAAGGAUAAUTT-3′ (GenePharma, Shanghai, China).

Statistical Analysis. All data were expressed as mean ± S.E.M. Statistical analysis was performed using one-way ANOVA followed by Newman-Keuls test or using two-way ANOVA (myocardial blood flow (MBF), left ventricular (LV) developed pressure (max), LV developed pressure (min) and LV systolic pressure) followed by Bonferroni for multiple comparisons. A p value less than 0.05 was considered to be statistically significant.

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**Acknowledgments**

This work was supported by the National Natural Science Foundation of China [81273637] to Jing-Yan Han.
Author Contributions
X.Y.Y. designed the experiments, acquired most data in all Figures and wrote the manuscript. K. H. acquired some supplementary results and revised the manuscript. C.S.P. participated in the design and interpretation of western and ELISA experiments. Q.L., S.W.H. and Y. J. participated in acquisition and interpretation of proteomics. Y.Y.L., L.Y., X.W.M. and D.D.H. participated in animal experiments. X.H.W. participated in histological experiment. B.H.H. and X. C. participated in the design and acquisition of electron microscope. L.J.W. participated in design and interpretation of SPR. G.C.W. provide the two modified DLA analogues. J.Y.F. and T.P.F. revised the manuscript. J.Y.H. designed, revised the manuscript, interpreted data and finally approved the submission of this manuscript.

Additional Information
Supplementary information accompanies this paper at http://www.nature.com/srep

Competing financial interests: The authors declare no competing financial interests.

How to cite this article: Yang, X.-Y. et al. 3, 4-dihydroxyl-phenyl lactic acid restores NADH dehydrogenase 1 α subunit 10 to ameliorate cardiac reperfusion injury. Sci. Rep. 5, 10739; doi: 10.1038/srep10739 (2015).