Early Donepezil Monotherapy or Combination With Metoprolol Significantly Prevents Subsequent Chronic Heart Failure in Rats With Reperfused Myocardial Infarction

Meihua Li (limehua@ncvc.go.p)
National Cerebral and Cardiovascular Center  https://orcid.org/0000-0003-1184-5482

Can Zheng
National Cerebral and Cardiovascular Center

Kawada Toru
National Cerebral and Cardiovascular Center

Kazunori Uemura
National Cerebral and Cardiovascular Center: Kokuritsu Junkankibyo Kenkyu Center

Inagaki Masashi
National Cerebral and Cardiovascular Center

Takuya Nishikawa
National Cerebral and Cardiovascular Center

Keita Saku
National Cerebral and Cardiovascular Center

Masaru Sugimachi
National Cerebral and Cardiovascular Center

Routine Article

Keywords: chronic heart failure, donepezil, metoprolol, myocardial salvage, infarct size, reperfused myocardial infarction

DOI: https://doi.org/10.21203/rs.3.rs-175205/v1

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Abstract

Purpose

Acute myocardial infarction (MI) remains a leading cause of chronic heart failure, and MI size is a strong predictor of prognosis. Clinical guidelines recommend that oral β-blockers be initiated early after reperfused MI (RMI). Here, we compared the effects of donepezil, metoprolol, and their combination on infarct size and progression of cardiac remodeling in rats with RMI.

Methods

RMI ($n = 103$) was induced in 8-week-old male rats by occluding the left coronary artery (30 min), followed by reperfusion. The animals were randomly assigned to untreated (UT, $n = 16$), donepezil-treated (DT, 5 mg/kg·day, $n = 21$), metoprolol-treated (MT, 70 mg/kg·day, $n = 23$), and combination of donepezil and metoprolol (DMT, $n = 18$) groups.

Results

On day 8 after surgery, rats in the DT and DMT groups exhibited significantly improved myocardial salvage, owing to the prevention of apoptosis and suppression of macrophage infiltration, compared with rats in the UT group. After the 10-week treatment, the rats in the DT and DMT groups exhibited decreased heart rate, reduced MI size, attenuated cardiac hypertrophy and cardiac dysfunction, and decreased plasma levels of brain natriuretic peptide, norepinephrine, and high-sensitivity C-reactive protein. However, no cardioprotective effects were observed among rats in the MT group.

Conclusions

Donepezil monotherapy or in combination with metoprolol significantly improved myocardial salvage, reduced MI size, limited the progression of cardiac remodeling, and prevented subsequent chronic heart failure in rats with RMI. Thus, donepezil monotherapy or combination with metoprolol may be applied as pharmacotherapy post-RMI.

Introduction

Timely reperfusion via primary percutaneous coronary intervention (PCI) is the best therapeutic strategy for patients with acute myocardial infarction (AMI), and its widespread implementation has significantly reduced patient mortality [1]. Nevertheless, many individuals with AMI develop cardiac remodeling and chronic heart failure (CHF) because of the loss of viable myocardium, even after reperfusion therapy [2–4][2–4]. Therefore, it is necessary to salvage more myocardium during the management of AMI to prevent subsequent CHF.

Sympathetic overactivity is involved in the pathology of CHF. Clinical practice guidelines recommend early (< 24 h) oral administration of β-blockers, such as metoprolol [5], after coronary reperfusion in
patients with AMI. However, oral metoprolol administration, which is initiated early post-reperfusion, does not improve myocardial salvage in patients with AMI [6]. Several experimental studies and clinical trials have reported controversial effects of early intravenous metoprolol administration before coronary reperfusion on myocardial salvage, MI size, and cardiac remodeling, and improvements in endpoints of efficacy were not confirmed in a larger population [7–10]. Decreased parasympathetic activity is another independent risk factor following AMI [11, 12]; however, few studies have investigated treatment alternatives to combat parasympathetic dysfunction.

Parasympathetic activation via electrical vagal nerve stimulation prevents cardiac remodeling and improves the long-term survival of rats with CHF [13]. Moreover, early vagal nerve stimulation attenuates cardiac remodeling after reperfused MI (RMI) in rabbits [14]. Recently, we proposed a novel pharmacotherapeutic approach using donepezil, an acetylcholinesterase inhibitor, to improve parasympathetic function. Oral administration of donepezil, as monotherapy or in combination with losartan, prevents the progression of cardiac remodeling and improves the long-term prognosis in rats with CHF with permanent MI [15, 16]. Thus, adding donepezil to conventional β-blocker therapy may improve the autonomic balance and prevent CHF.

Here, we examined whether early administration of donepezil, metoprolol, or their combination in a rat model of RMI affected myocardial salvage, infarct size, cardiac remodeling, cardiac dysfunction, and the development of CHF. Our findings provide a rationale for early administration of donepezil as an adjunct therapy to PCI, thereby improving the long-term prognosis of patients with AMI.

**Methods**

Animals were housed and studied in accordance with Directive 2010/63/EU of the European Parliament and the Guiding Principles for the Care and Use of Animals in the Field of Physiological Science. The investigation conformed to the NIH Guide for the Care and Use of Laboratory Animals (NIH Publication No. 85–23, revised 1985). All protocols were reviewed and approved by the Animal Subject Committee of the National Cerebral and Cardiovascular Center (Osaka, Japan). The experiment comprised two studies: myocardial salvage and cardiac remodeling. During experiments, the animals were anesthetized with halothane (3% at induction, 1.2% during surgery, and 0.6% during data recording), and the body temperature was maintained at 37°C. For details, see the Methods section of the online Supplementary Material.

**RMI model**

Eight-week-old male Sprague-Dawley rats (SLC, Inc., Hamamatsu, Japan) were used (n = 141). After inducing anaesthesia, the heart was exposed, and the pericardium was opened via a left lateral thoracotomy at the second intercostal space. A 5–0 prolene suture was placed around the left ventricular proximal coronary artery (LCA) to create a reversible snare. All rats underwent a total of 30 minutes of...
LCA occlusion, followed by reperfusion and closed the chest. The loosened snare was kept in the chest for identifying the ischaemic regions later.

**Experimental protocol**

Shortly after induction of RMI, 23 rats were used for the myocardial salvage study, and 80 rats were used for the cardiac remodeling study (Fig. 1A). All drug treatments were initiated 1 h after recovery from anaesthesia. Rats with RMI were randomly divided into the untreated (UT), donepezil-treated (DT), metoprolol-treated (MT), and donepezil plus metoprolol-treated (DMT) groups. For the DT and DMT groups, donepezil was dissolved in drinking water to provide a dose of 5 mg/kg-day. For the MT and DMT groups, metoprolol was dissolved in drinking water at a dose of 70 mg/kg-day. These dosages were selected as previously described [17, 15].

**Myocardial salvage study**

After 1 week of treatment, rats with RMI (Fig. 1B) were analysed as described below.

**Determination of neutrophil infiltration**

The middle of the short-axis portion of the left ventricle (LV) was embedded in paraffin, sectioned transversely to 4-µm thickness, and stained with Masson's trichrome. Neutrophils were counted using a light microscope (BZ-9000; KEYENCE, Osaka Japan) coupled to an image-analysis software at 40× magnification in six fields, which were randomly selected from the ischaemic/reperfusion (I/R) region per rat.

**Analysis of myocardial apoptosis**

Myocardial apoptosis in the I/R region was analysed. Apoptotic cells were detected using terminal deoxynucleotidyl transferase dUTP nick end labelling (TUNEL) staining [18]. Biventricular sections were deparaffinized and incubated with the TUNEL reagent and rabbit anti-connexin 43 antibody, followed by incubation with Alexa Fluor 633-conjugated goat anti-rabbit IgG antibody. The fluorescence of Alexa Fluor 488 (TUNEL), Alexa Fluor 633 (connexin 43), and Alexa Fluor 350 (nuclei) was observed under a fluorescence microscope. Apoptotic myocardial cells were counted at 20× magnification. Data obtained from six fields were averaged and expressed as the number of apoptotic cells.

**Examination for macrophage infiltration**

Myocardial salvage associated with myocardial inflammation in the I/R region, which was assessed by observing for macrophage infiltration. Tissue sections were prepared to examine macrophage infiltration and incubated with mouse anti-CD68 and rabbit anti-human von Willebrand factor (vWF) polyclonal antibodies. Sections were then incubated with Alexa Fluor 488-conjugated goat anti-mouse IgG and Alexa Fluor 633-conjugated goat anti-rabbit IgG antibodies. The fluorescence of Alexa Fluor 488 (CD68), Alexa Fluor 633 (vWF), and Alexa Fluor 350 (nuclei) was observed using a fluorescence microscope at 40× magnification. To distinguish between classically activated pro-inflammatory type 1 macrophages (M1)
and alternatively activated anti-inflammatory type 2 macrophages (M2), the following were stained together: (1) CD68, CD80, and DAPI for M1; or (2) CD68, CD163, and DAPI for M2 [19]. Data obtained from six fields were averaged and expressed as the percentage area of macrophage infiltration.

**Cardiac remodeling study**

**Telemetric measurements of blood pressure (BP) and heart rate (HR)**

Rats were anesthetized and ventilated through an endotracheal cannula during the implantation procedure. Rats were implanted with BP transmitters (TA11PA-C40; DSI, St. Paul, MN, USA) for real-time monitoring of the hemodynamics of conscious rats. BP transmitter catheters were inserted into the abdominal aorta. Data on BP and HR of freely moving animals were recorded continuously. The recording was sampled at 500 Hz. One week after implantation of BP transmitters, RMI was induced, and 80 surviving rats were randomly assigned to the UT, DT, MT, and DMT groups. After 10 weeks of treatment, the surviving rats (Fig. 1C) were analysed.

**Hemodynamic measurements and assessment of cardiac remodeling under anaesthesia**

Ten weeks after RMI, the hemodynamics and heart weights of rats with CHF were determined. LV and arterial pressures were measured with a 2-Fr catheter-tip micromanometer (SPC-320; Millar Instruments), and aortic flow was measured with a flow probe (Transonic Systems Inc.; T-206 flow probe #2.5 ss66). Right atrial pressure (RAP) was measured using a fluid-filled pressure sensor. All signals were digitized at 500 Hz for 5 min. After hemodynamic measurements, 2 mL blood was sampled and transferred into an EDTA-2Na tube. Neurohumoral and cytokine assays were performed with the plasma, which was obtained after centrifugation of the blood (3000 × g, 20 min) at 4°C.

**Neurohumoral and cytokine assays**

Plasma concentrations of catecholamines were measured using high-performance liquid chromatography with electrochemical detection after alumina adsorption [15]. Plasma levels of brain natriuretic peptide (BNP) and high-sensitivity C-reactive protein (hs-CRP) were determined using enzyme-linked immunosorbent assay kits, per the manufacturer's instructions.

**Histological assessments of risk area, infarct size, cardiac fibrosis, and myocyte cross-sectional area**

Rats were euthanized with an overdose of pentobarbital sodium (100 mg/kg, i.v.) after hemodynamic measurements and blood sampling during anaesthesia. Hearts were harvested and mounted on a modified [19, 18] Langendorff apparatus. The perfusate was infused with Pigment Blue Ink (PLATINUM CO., LTD., Japan) to identify the ischaemic regions. Once the dye had stained the non-ischaemic regions, biventricular weights were determined, and the hearts were sectioned. Three portions (1-mm thick) from
the apex to the base were selected and embedded in paraffin. Sections (4-µm thick) were cut and stained with Masson's trichrome. Histological images were obtained with a microscope. The area at risk was calculated as the percentage of the unstained I/R region relative to the whole LV area, which was averaged for the three portions. The infarcted area was calculated as the percentage of the area of scar tissues relative to the whole LV area, which was averaged for the three portions. MI size was defined as the percentage of the infarcted area relative to the area at risk. The extent of cardiac fibrosis was evaluated using a light microscope at 20× magnification, and myocyte cross-sectional area (papillary cardiomyocytes) was evaluated at 40× magnification. The area of cardiac fibrosis in each heart was calculated as the ratio of the blue area to the total tissue area in eight fields of the non-infarcted area.

**Immunohistochemical examination and analysis of microvessel density**

Biventricular sections were pretreated as described in the myocardial salvage study. Sections were incubated with a rabbit anti-human vWF polyclonal antibody and then with Alexa Fluor 633-conjugated goat anti-rabbit IgG antibody for analysis of microvessel density. Capillary vessels in the peri-infarct area and septum were counted as fluorescent regions using Alexa Fluor 633 (vWF) and a laser-scanning microscope at 20× magnification. Data obtained from 2–5 fields were averaged and expressed as the number of capillary vessels.

**Statistical analysis**

All statistical analyses were performed using GraphPad Prism 7 (GraphPad Software, San Diego, CA, USA). All data were expressed as means ± standard errors of the means (SEMs). Long-term-recorded data on HR and mean BP (MBP) before and during treatment within each group were compared using repeated measures one-way analysis of variance (ANOVA), followed by Dunnett's test. For data obtained from the hemodynamic and remodeling study, differences between groups were tested using one-way ANOVA, followed by Dunnett's multiple comparisons test. Data on the myocardial salvage study, cardiac fibrosis, MI size, microvessel density, neurohumoral, and cytokine assays were analysed using nonparametric Kruskal-Wallis tests, followed by Dunn's tests for all pairwise comparisons. Differences were considered significant at the level of $P < 0.05$. 

**Results**

**Immune cell infiltration into cardiac cells and myocardial apoptosis**

Figure 2 shows representative images of Masson's trichrome (Fig. 2A, B, S1) and TUNEL-Cox43 (Fig. 2D) staining of hearts on day 8 after RMI. Considerable neutrophil infiltration into the myocardium of RMI tissues was observed in the UT and MT groups, with significant reductions in the DT and DMT groups (Fig. 2C). Apoptotic cell counts were significantly lower in the DT and DMT groups than in the UT and MT groups (Fig. 2E), with strong connexin43 signals in the DT and DMT groups.
Immunohistochemical staining of CD68+ cells (all macrophages), CD68+-vWF+ cells (microvessels), CD68+-CD80+ (M1) cells, and CD68+-CD163+ (M2) cells in RMI tissues revealed decreased total macrophage infiltration, decreased M1 cells, and increased M2 cells and microvessels in the I/R region in the DT and DMT groups, compared with that in the UT and MT groups (Fig. 3A–H).

**Telemetric hemodynamic measurements in conscious rats with RMI-induced CHF**

In the cardiac remodeling study, the weekly average HR in the UT group increased to 413 bpm at week 1 during pretreatment and then decreased (Fig. 4A). Compared with the UT group, the weekly average HR in the other groups was significantly reduced. This reduced HR was prominent in the DMT group. The difference in HR between the DMT and UT groups was significant and approximately 40 bpm at week 4 of treatment. Figure 4B shows changes in daily average HR during the first 7 days. The HR increased, reached a maximum (425 bpm) on day 3, and then gradually decreased in the UT group. The difference in HR between the DMT and UT groups was significant and approximately 60 bpm on day 7. Similarly, the HR in the DT and MT groups was decreased significantly compared with that in the UT group. The weekly MBP in the DT, MT, and DMT groups increased from that at week 0, but did not significantly change in the UT group (*Table S1*). The differences in MBP reached approximately 10 mmHg at the end of week 4.

**Hemodynamics under anaesthesia, cardiac remodeling, and cardiac function**

The parameters of hemodynamics, cardiac remodeling, and cardiac function after 10 weeks of treatment are shown in Table 1. Body weights of the rats in the DT, MT, and DMT groups were higher than those in the UT group. Rats in the DT and DMT groups had lower heart weights, higher cardiac index (CI), lower LV end-diastolic pressure (LVEDP), higher maximum dp/dt LV pressure (LV dp/dt\textsubscript{max}), and minimum dp/dt LV pressure (LV dp/dt\textsubscript{min}) than the rats in the UT group.
Table 1
Hemodynamic, cardiac remodeling, and plasma neurohumoral and inflammatory parameters in rats with RMI-induced chronic heart failure (CHF) after 10 weeks of treatment

|                      | UT group (n = 12) | DT group (n = 15) | MT group (n = 18) | DMT group (n = 14) |
|----------------------|-------------------|-------------------|-------------------|--------------------|
| BW, g                | 495 ± 7           | 523 ± 7           | 525 ± 10          | 545 ± 9**          |
| HW, g/kg             | 2.73 ± 0.13       | 2.39 ± 0.05*††    | 2.73 ± 0.06       | 2.41 ± 0.03*††     |
| Infarcted area, %    | 20.1 ± 1.2        | 12.5 ± 1.5**††    | 21.1 ± 1.1        | 14.3 ± 1.7*††      |
| MBP, mmHg            | 96 ± 3            | 98 ± 3            | 96 ± 3            | 107 ± 3            |
| HR, bpm              | 302 ± 7           | 312 ± 6           | 304 ± 4           | 317 ± 7            |
| CI, mL/min/kg        | 98 ± 6            | 120 ± 4*††        | 91 ± 4            | 120 ± 4*††         |
| LV dp/dt_{max}, mmHg/s | 4318 ± 127       | 4853 ± 173*       | 4387 ± 180        | 4950 ± 120*†       |
| LV dp/dt_{min}, mmHg/s | 3493 ± 147       | 4022 ± 154*††     | 3511 ± 166        | 4277 ± 113*††     |
| LVEDP, mmHg          | 23 ± 3            | 14 ± 2*           | 21 ± 2            | 15 ± 1*            |
| RAP, mmHg            | 2.9 ± 0.3         | 3.1 ± 0.4         | 3.4 ± 0.3         | 2.6 ± 0.5          |
| Plasma:              |                   |                   |                   |                    |
| NE, pg/mL            | 392 ± 96          | 155 ± 27*         | 282 ± 58          | 111 ± 18**†        |
| Epi, pg/mL           | 430 ± 85          | 112 ± 55**††      | 381 ± 92          | 119 ± 22**†        |
| BNP, pg/mL           | 553 ± 42          | 370 ± 16**††      | 482 ± 20          | 417 ± 8**          |
| hs-CRP, ng/mL        | 825 ± 67          | 262 ± 61**        | 415 ± 38*         | 201 ± 12**††       |

BW, body weight; HW, biventricular weight normalized by body weight; MBP, mean arterial blood pressure; HR, heart rate; CI, cardiac index; LV dp/dt_{max}, maximum dp/dt of the left ventricular pressure; LV dp/dt_{min}, minimum dp/dt of the left ventricular pressure; LVEDP, left ventricular end-diastolic pressure; RAP, right atrial pressure. NE, norepinephrine; Epi, epinephrine; BNP, brain natriuretic peptide; hs-CRP, high-sensitivity C-reactive protein. Values are expressed as means ± SEMs. *P < 0.05; **P < 0.01 versus UT group, † P < 0.05; †† P < 0.01 versus MT group. For data from the hemodynamic and remodeling studies under anaesthesia, differences between the UT, DT, MT, and DMT groups were tested using one-way ANOVA, followed by Tukey's multiple comparison tests. For data on plasma neurohumoral and inflammatory parameters, differences between UT, DT, MT, and DMT groups were tested using nonparametric Kruskal-Wallis tests followed by Dunn's tests for all pairwise comparisons. UT, untreated; DT, donepezil treatment; MT, metoprolol treatment; DMT, donepezil plus metoprolol treatment.

Histological assessments of risk area, MI size, cardiac fibrosis, and myocyte cross-sectional area
Among rats in the DT and DMT groups, histological analyses revealed the association of preserved cardiac function with histological changes (Fig. 5A–C, S2). Rats in the DT and DMT groups showed a significantly reduced MI size compared with that in the UT group. Similarly, among the rats in the DT and DMT groups, cardiac fibrosis was significantly suppressed, and cardiac myocyte hypertrophy was markedly attenuated as a direct consequence of the reduced myocyte cross-sectional area (Fig. 5D–G). These beneficial effects were not observed among the rats in the MT group.

**Analysis of microvessel density**

Immunohistochemical analysis of vWF in the peri-infarct area and septum revealed that neovascularization in the DT and DMT groups was more pronounced than that in the UT group (Fig. 5H). Quantitative analysis confirmed a significantly higher capillary density in the peri-infarct and septum regions in the DT and DMT groups than in the UT group. Metoprolol monotherapy did not promote angiogenesis in RMI-induced rats with CHF (Fig. 5I, J).

**Neurohumoral and cytokine levels**

Table 1 shows the plasma levels of neurohumoral and inflammatory markers after 10 weeks of treatment. Rats in the DT and DMT groups showed lower plasma levels of norepinephrine, epinephrine, and BNP and reduced hs-CRP levels compared with rats in the UT group. Metoprolol monotherapy reduced plasma hs-CRP levels compared with those in the UT group.

**Discussion**

We investigated the effects of early administration of donepezil, metoprolol, and their combination on cardiac function in a rat model of RMI. We showed that early administration of donepezil (with or without metoprolol) facilitated myocardial salvage, prevented HR increase and preserved MBP, reduced MI size, prevented the progression of cardiac remodeling and dysfunction, suppressed catecholamine and BNP plasma levels, promoted angiogenesis, and suppressed inflammation in the heart. Thus, early administration of donepezil (with or without metoprolol) prevented structural changes in the heart after RMI, probably via activation of the vagal nerve. Local anti-inflammatory effects on the heart early in treatment may contribute to myocardial salvage. Early administration of donepezil may be effective for post-RMI therapy.

**Early donepezil treatment (with or without metoprolol) affected myocardial salvage post-RMI via an anti-inflammatory mechanism**

Donepezil (with or without metoprolol), initiated early post-RMI, preserved myocardial salvage by suppressing neutrophil infiltration and myocardial apoptosis in the I/R region, 1 week after RMI. These treatments also attenuated total macrophage infiltration and suppressed local inflammation by reducing
the production of pro-inflammatory M1 cells and promoting the production of anti-inflammatory M2 cells. The M1/M2 ratio was reduced in the I/R region. Suppression of local inflammation reduced infarct size, limited cardiac remodeling, preserved cardiac function and prevented subsequent progression to CHF 10 weeks post-RMI. When uncontrolled, these immune responses, which are related to the structural changes after RMI, induce cardiac remodeling, leading to CHF. Specifically, MI size is a strong predictor of cardiac remodeling, dysfunction, and cardiovascular events, which subsequently progress to CHF [20]. Furthermore, myocardial salvage is associated with the effects of myocardial inflammation, including myocardial apoptosis and macrophage/neutrophil infiltration in the I/R region, early post-RMI [6]. These observations are consistent with our previous reports showing anti-inflammatory effects of donepezil in rats with CHF and permanent MI [15, 21].

**Early donepezil, metoprolol, and combined donepezil-metoprolol therapy reduced HR and affected MBP preservation in RMI-induced CHF**

We used a telemetric system to monitor arterial MBP and HR in conscious, unstressed rats. Donepezil treatment (from week 2), metoprolol treatment, or their combination (from week 1) reduced the weekly average HR compared with that in the UT group. In particular, combined treatment significantly decreased the immediate increase in HR from day 2 post-RMI, and metoprolol treatment decreased the increase in HR from day 3 post-RMI. Furthermore, weekly HR in the DMT group nearly reached that of healthy animals at week 4 post-RMI [13]. The observed reduction in HR may have resulted from a decreased sympathetic drive and increased efferent discharges of the parasympathetic system [12]. Although we did not directly evaluate sympathetic nerve activity, low plasma levels of norepinephrine and epinephrine in the DT and DMT groups may reflect decreased central sympathetic outflow [22]. The bradycardia observed in the DT and DMT groups may play important roles in preventing cardiac dysfunction. Additionally, the effects of bradycardia were observed in the MT group. Metoprolol is a β-blocker, which induces bradycardia, and is important for the treatment of patients with CHF. However, because the sinus node and cardiac myocytes express β-adrenergic receptors, β-blockers decrease HR and suppress myocardial contractility. Hence, considering the negative inotropic effect, β-blockers may not be suitable for patients with poor hemodynamics or decompensation with pre-existing myocardial dysfunction because the maintenance of cardiac output in such patients partly depends on an increased sympathetic drive. Hence, although metoprolol treatment led to a long-term decrease in HR, it did not significantly improve LV dp/dt\text{\textsubscript{max}} or LV dp/dt\text{\textsubscript{min}}, resulting in low CI and high LVEDP. The plasma levels of catecholamines were also not significantly affected in the MT group compared with the levels in the UT group. Although metoprolol treatment blocks peripheral sympathetic effects, it did not decrease the central sympathetic outflow in CHF. Vasoconstriction could occur via unblocked α-adrenergic receptors, which may have contributed to the increased MBP in the MT and DMT groups. However, donepezil treatment also increased MBP over the baseline (week 0) values. Because the vagus nerve does not directly affect ventricular contractility, increases synaptic acetylcholine levels, and reduces HR, the negative effects on the pumping function of the heart are lower with donepezil than with metoprolol. Prevention of cardiac dysfunction may also have contributed to the maintenance of MBP, resulting in
arterial baroreflex-mediated inhibition of sympathetic nerve activity and HR reduction [23]. A prolonged cardiac cycle is beneficial for the enhancement and maintenance of cardiac function by decreasing myocardial oxygen consumption, increasing coronary blood flow, and increasing ventricular filling time [24]. Since acetylcholine antagonizes the effects of β-adrenergic stimulation [25], donepezil may be either an alternative to β-blockers or adding to the conventional β-blockers therapy in patients with CHF post-RMI.

**Early donepezil (with or without metoprolol) affected cardiac remodeling and dysfunction**

Infarct size is a strong predictor of cardiovascular events. We showed that donepezil treatment, as monotherapy or in combination with metoprolol, reduced infarct size, prevented cardiac fibrosis and cardiomyocyte hypertrophy, and increased microvessel density. These treatments prevented the progression of cardiac remodeling and dysfunction compared with that in untreated rats. Plasma catecholamine levels were also significantly reduced in the DT and DMT groups, consistent with our previous studies [15, 16]. Furthermore, plasma BNP levels in the DT and DMT groups were significantly reduced compared with those in the UT group, which may have resulted from the augmented vagal tone and decreased central sympathetic outflow [22]. These observations showed that early administration of donepezil (with or without metoprolol) after RMI is beneficial mainly via local anti-inflammatory effects and prevention of myocardial apoptosis, showing myocardial salvage in the acute phase. Metoprolol monotherapy did not exert these beneficial effects, but reduced hs-CRP levels and HR and preserved MBP. The reduced hs-CRP levels in the MT group may indicate an indirect reduction in the inflammatory response of the tissues, which is associated with reduced shear stress due to long-term reduction of HR.

**Probable mechanisms of the effects of early administration of donepezil (with or without metoprolol)**

In the DT and DMT groups, suppression of local inflammation 1 week after RMI may be associated with decreased plasma hs-CRP levels and increased microvessel density in the peri-infarct and remote regions after 10 weeks of treatment. This leads to more pronounced myocardial salvage than in the UT group by preventing enlargement of MI size and suppressing cardiac fibrosis and cardiomyocyte hypertrophy. Although PCI entails second-generation drug-eluting stents and reduces post-AMI care, the incidence of post-AMI CHF remains high; every 5% increase in infarct size is associated with a 20% increase in 1-year hospitalisation for CHF and 1-year mortality [26–28]. Here, we showed that infarct size was decreased by 35% and 31% in the DT and DMT groups, respectively, compared with that in the untreated group. Metoprolol treatment at 2 weeks after permanent MI improved cardiac function [17]; however, in this study, early metoprolol monotherapy did not exert beneficial effects, excluding the reductions in hs-CRP levels and HR and preservation of MBP. The effects of DMT were additive. Thus, early donepezil treatment, either monotherapy or in combination with metoprolol, may be a novel pharmacotherapy, as an adjunct therapy to PCI, to improve the long-term prognosis in patients with AMI.

**Limitations**
In this study, we used an RMI-induced CHF rat model to evaluate the precarious clinical condition of hospitalised patients with AMI. However, the experimental animals were young, with possible preservation of autonomic function, and were more responsive to various therapeutic interventions. However, many patients with AMI are middle- to old-aged and may have limited capacity for response to the testing of this novel treatment. Furthermore, because clinical trials usually involve patients with various backgrounds of pharmacological treatment, it is difficult to determine the efficacy of monotherapy in that population. This may be an important factor to be considered when translating the outcomes of this basic study to clinical practice.

Conclusions

Donepezil monotherapy or in combination with metoprolol significantly improved myocardial salvage, reduced MI size, limited the progression of cardiac remodeling and dysfunction, and prevented subsequent CHF. This suggests that donepezil, with or without metoprolol, may be a potential candidate for post-RMI therapy.

Declarations

Funding

This work was partly supported by JSPS KAKENHI (grant numbers 17K09544, 18K08091)

Conflicts of interest/Competing interests

None declared.

Ethics approval

The care of animals and all animal experiments were performed in strict accordance with the guiding principles of the Physiological Society of Japan and the Guide for the Care and Use of Laboratory Animals published by the US National Institutes of Health (NIH Publications No 85-23, revised 1996). All protocols were reviewed and approved by the Animal Subject Committee in the National Cerebral and Cardiovascular Center.

Consent to participate

Not applicable

Consent for publication

Not applicable

Availability of data and material
The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

**Code availability**

None

**Author contributions**

ML and CZ designed the study. ML and CZ performed the measurements and statistical analyses and drafted the manuscript. TK, KU, MI, TN, KS and MS joined in interpreting the data. ML and CZ wrote and edited, TK and KS reviewed the manuscript. All authors have read and approved the final manuscript.

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