Overexpression of *miR-1* in the heart attenuates hippocampal synaptic vesicle exocytosis by the posttranscriptional regulation of SNAP-25 through the transportation of exosomes

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Abstract

**Background:** The link between cardiac diseases and cognitive deterioration has been accepted from the concept of "cardiogenic dementia", which was proposed in the late 1970s. However, the molecular mechanism is unclarified.

**Methods:** The two animal models used in this study were cardiac-specific overexpression of microRNA-1-2 transgenic (Tg) mice and a myocardial infarction mouse model generated by left coronary artery ligation (LCA). First, we observed the *miR-1* (*miR-1*) level and synaptic vesicles (SV) distribution in the hippocampus using in situ hybridization and transmission electron microscopy (TEM) and evaluated the expression of vesicle exocytosis related proteins by western blotting. Second, we used dual luciferase reporter assay as well as antagonist and miRNA-masking techniques to identify the posttranscriptional regulatory effect of *miR-1* on the *Snap25* gene. Third, FM1–43 staining was performed to investigate the effect of *miR-1* on synaptic vesicle exocytosis. Lastly, we used GW4869 to inhibit the biogenesis and secretion of exosomes to determine the transportation effect of exosomes for *miR-1* from the heart to the brain.

**Results:** Compared with the levels in age-matched WT mice, *miR-1* levels were increased in both the hearts and hippocampi of Tg mice, accompanied by the redistribution of SVs and the reduction in SV exocytosis-related protein SNAP-25 expression. In vitro studies showed that SNAP-25 protein expression was down- or upregulated by *miR-1* overexpression or inhibition, respectively, however, unchanged by miRNA-masking the 3'UTR of the *Snap25* gene. SV exocytosis was inhibited by *miR-1* overexpression, which could be prevented by co-transfection with an anti-*miR-1* oligonucleotide fragment (AMO-1). The knockdown of *miR-1* by hippocampal stereotaxic injection of AMO-1 carried by a lentivirus vector (lenti-pre-AMO-1) led to the upregulation of SNAP-25 expression and prevented SV concentration in the synapses in the hippocampi of Tg mice. The application of GW4869 significantly reversed the increased *miR-1* level in the blood and hippocampi as well as reduced the SNAP-25 protein levels in the hippocampi of both Tg and LCA mice.

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Conclusion: The overexpression of miR-1 in the heart attenuated SV exocytosis in the hippocampus by posttranscriptionally regulating SNAP-25 through the transportation of exosomes. This study contributes to the understanding of the relationship between cardiovascular disease and brain dysfunction.

Keywords: Heart, Hippocampus, Synaptic vesicle exocytosis, microRNA-1, SNAP-25

Background
The primary function of microRNAs (miRNAs) is post-transcriptionally regulating target protein expression by binding to their mRNA recognition sequences [1]. However, with the increasing number of miRNA studies, miRNAs were found to be stable in various body fluids (i.e., serum, plasma, saliva, urine, breast milk, and tears), and showed potential as biomarkers for various diseases [2–9]. More excitingly, miRNAs are also mediate trans-regulation between different cell types after these molecules are secreted into the interstitial space from donor cells and accepted by close recipient cells with functionally targeting capabilities through the actions of various transporters [10–13]; miRNAs even mediate long-distance metabolic regulation from adipose to the liver [14] and in cancer metastasis [15, 16]. However, whether miRNAs could mediate long-distance heart-brain communication and act as a link between heart disease and brain dysfunction is largely unknown, which would be very interesting.

MicroRNA-1 (miR-1) is a muscle-enriched miRNA that presents a hundred-fold lower level in the brain than in the heart [17, 18]. Interestingly, by bioinformatics analysis, we found that as a muscle-enriched microRNA, 57% of the targets of miR-1 were distributed in the brain and only 5% of the targets were distributed in the heart of adult mice (Fig. 1). Our previous study demonstrated

![Fig. 1](image_url) Distribution of miR-1 targets in different tissues. By uploading the 398 target genes onto the David bioinformatics web site, we obtained the tissue distribution of 358 miR-1 targets. All of the miR-1 targets were classified into 3 parts: targets in fetal tissue, targets in mature tissue and targets associated with tumourigenesis. In each part, we analysed the percentage of miR-1 targets in each kind of tissue. For example, in mature tissue, 57% of the targets are distributed in brain tissue and 5% are located in the heart (red arrow).
the behavioural abnormalities in a cardiac-specific microRNA-1-2 (miR-1-2) overexpression transgenic (Tg) mouse model, which was associated with the downregulation of BDNF expression in the hippocampus [18]. Furthermore, a very recent study demonstrated that the overexpression of miR-1 in the heart could induce neuronal microtubule damage [19]. These studies revealed the potential of cardiac-originated miRNAs to regulate brain function. Indeed, clarifying the underlying biological significance of heart-brain communication mediated by miRNAs would provide a new insight into association to the prevention of brain dysfunction.

Synaptic pathology is one of the prominent features of Alzheimer’s disease (AD) or vascular dementia (VaD) [20]. Considering the relationship between cardiovascular diseases and dementia [17, 21–25], we hypothesized that the overexpression of miR-1 in the heart might be involved in heart disease-induced abnormal synaptic plasticity. The study will provide evidence to reveal a novel insight into the molecular mechanism of the heart-to-brain connection and disclose a potential communication between cardiac disease and brain disorder.

**Methods**

**Animals**

Adult male C57BL/6 mice and cardiac-specific miR-1 overexpression transgenic (Tg) mice (6–7 months) were housed under a controlled temperature of 23 ± 1 °C and humidity of 55 ± 5%. The animals were maintained on a 12 h artificial dark-light cycle (lights on at 07:00 A.M.) with food (regular chow) and water ad libitum. All animal procedures were approved by the ethics committee with food (regular chow) and water ad libitum. All animal procedures were approved by the ethics committee of Harbin Medical University and carried out in accordance with the European Communities Council Directive of November 24, 1986 (86/609/EEC).

**miR-1 targets and bioinformatic analysis**

We acquired mouse miRNA target genes from 7 miRNA target predicting tools, including PicTar, RNAhybrid, DIANA-microT, RNA22, miRBase Targets, miRanda and TargetScan. To improve the reliability of the predicted miRNA regulations, we only extracted the regulations that were predicted by at least two tools. We obtained 398 target genes. Finally, by uploading the 398 target genes onto the David bioinformatics web site, we obtained the tissue distribution of 358 miR-1 targets. After double verification by literature citation, the percentages of the target gene locations were calculated.

**Generation of miR-1 transgenic (Tg) mice**

**miR-1** Tg mice were generated as previously described [26]. Sexually immature female C57BL/6 mice (4–5 weeks of age) were used to obtain sufficient quantity of eggs (> 250) for microinjection. The Tg mice used in this study were 5th generation or later. Five- and six-month-old mice (5 M and 6 M) were used in this experiment.

**In situ hybridization (ISH)**

First, 10 μm paraffin sections were prepared from the hippocampi and hearts of mice. After deparaffinization with xylene and ethanol, the sections were washed 3 times with a phosphate buffer solution (PBS) buffer for 5 min each. Then, proteinase K (40 μg/mL) was applied to deproteinate the sections for 20 min at room temperature. Second, after re-fixing in 4% paraformaldehyde (PFA) for 10 min and acetylating in 1.15% triethanolamine and 0.25% acetic anhydride, the sections were prehybridized in hybridization buffer (50% formamide, 5x standard saline citrate (SSC), 0.1% Tween-20, 50 μg/mL heparin, 500 μg/mL yeast tRNA, and 9.2 mM citric acid for adjustment to pH 6) for 2 h at 37 °C. The digoxigenin (DIG)-labelled detection probes mmu-miR-1 (30 nM, No. 21378–15) and scramble-miR-1 (mis-miR-1, 30 nM, No.99004–15, Exiqon, Vedbaek, Denmark) were added to the slides and hybridized with the tissue overnight at 50 °C in a humidified chamber. The next day, the slides were washed twice with 5x SSC solution (20× SSC, 3 M NaCl, and 0.3 M sodium citrate) for 20 min each, and 3 times with 50% formamide at 37 °C for 20 min each and 5 times with TBS-T sequentially. After blocking for 1 h in blocking solution (2% goat serum, 2 mg/mL BSA, 0.1% Tween-20, and 1x PBS), the sections were incubated with anti-Digoxigenin-AP (1:1000, product No.11093274, Roche, Indianapolis, IN, USA) in blocking solution overnight at 4 °C. Finally, after washing 4 times with TBS containing 0.1% Tween-20, the slides were treated with developer buffer (0.1 M Tris, 0.1 M NaCl, 0.05 M MgCl2, 0.1%MTween, pH 9.5) for 20 min. Colour development was performed with 1-Step™ NBT/BCIP (product No.34042, Thermo Scientific, Rockford, USA). Images were captured using a microscope (AXIO, Zeiss, Germany).

**Mouse model of myocardial infarction**

The method was described in a previous study [27]. The male C57BL/6 wild-type mice were divided into two groups: the sham group and the myocardial infarction (MI) group. Briefly, the mice were anaesthetized with sodium pentobarbital (40 mg/kg, intraperitoneal) and placed in the supine position with the upper limbs taped to the table. A 1-cm incision was made along the left side of the sternum, and the muscle layers of the chest wall were bluntly dissected to avoid bleeding. The thorax was cut open at the point of the most pronounced cardiac pulsation, the left anterior descending coronary artery was ligated with a 7-0 silk suture and then the chest was closed. Ischaemia was confirmed by an elevated ST segment of the electrocardiogram.
Sham-operated control mice underwent the same surgical procedures except that the suture placed under the left coronary artery was not tied. All surgical procedures were performed under sterile conditions. After occlusion for 6 h, 1 d, 15 d and 30 d, hippocampi were collected for next experiments.

Evaluation of ultrastructure morphology by transmission electron microscopy

For transmission electron microscopy detection, the mice were anaesthetized with sodium pentobarbital (100 mg/kg). The hippocampi were removed and immersed in stationary liquid (pH 7.3) containing 3% glutaraldehyde in 0.1 mmol/L sodium phosphate buffer and 0.45 mmol/L CaCl₂. The tissue samples were then fixated in 2% osmic acid (OsO₄) in PBS with 1.5% potassium ferrocyanide. After dehydration with a concentration gradient of alcohol solutions, the tissues were embedded in Epon with propylene oxide as an intermediary solvent. Ultrathin sections were stained with uranyl acetate and lead citrate. The images were examined under a Hitachi H-7650 electron microscope (Hitachi, H-7650, Tokyo, Japan). To analyse vesicular distribution, 20 synapses without organelles near the presynaptic membrane were selected in each group (3 animals). The active zone was identified by a dark thickened presynaptic membrane with a visible synaptic cleft between the pre- and postsynaptic membranes. Both active zone length and the number of synaptic vesicles were measured using Image J software (National Institutes of Health, US). The total number of synaptic vesicles and vesicles located at 0 nm, 0–50 nm and 50–100 nm distances away from the active zone were counted in each selected pre-synapse [28].

Oligonucleotide synthesis

MiR-1 mimics and AMO-1 for rats were synthesized by Shanghai GenePharma Co., Ltd. (Shanghai, China). AMO-1 contains 2′-O-methyl modifications. In addition, scrambled RNA (mis-miR-1) was used as a negative control. Snap25-masking antisense oligodeoxynucleotides (ODNs) were synthesized by Shanghai Sangon Biotech Co., Ltd., China, and these ODNs complemented the position of 408–430 containing the binding sites of miR-1 located at position 413–420 in the 3′ UTR of Snap25. The nucleotides or deoxynucleotides at both ends of the antisense molecules were locked by a methylene bridge connecting the 2′-O and 4′-C atoms. The sequences of these synthesized oligonucleotides are shown in Table 1.

Construction of lentivirus vectors

To produce a miR-1 antisense inhibitor, two single-stranded DNA oligonucleotides were designed as follows: (1) pre-AMO-1 (“top strand” oligo: tgctg ATACATACT

Table 1 The sequences of the synthesized oligonucleotides

| Oligonucleotide | Sense | Antisense |
|----------------|-------|-----------|
| miR-1 mimics | 5′-UUGAAUGUAAAGAGUUGUACGU-3′ | 5′-AUACACACUUCUUUUAUCCAAU-3′ |
| AMO-1 | 5′-ACCUAUACAUUUUCUCACACAUACA-3′ | 5′-CUUCGCCACGGUGACGUAC-3′ |
| mis-miR-1 | 5′-UUCUGCCGACGGUGACGUAC-3′ | 5′-ACGUGACGUUGUGAGAAAU-3′ |
| Snap25-ODN | 5′GUAGCUCUGUCCGAUG-3′ | 5′GUAGCUCUGGUCCGAUGACAG-3′ |

TCTTTACATCCAGTTTTGCGCAGTACTGGATTGAGAAAGUGUAUGUGU-3′ and its complementary sequence (“bottom strand” oligo: cctgATACATACTTCTAC ATTCCATGTCAGTCCATTGCAAAACTGGAAATGTGAA AGAAGTATGTAT). (2) pre-mis-AMO-1 as a negative control (“top strand” oligo: tgctgAAATGTACTGGCGGGAGACGT TCGAGACGTCTTGGGCACTGACTGCTCCAGCAGTACATTTC) and its complementary sequence (“bottom strand” oligo: cctgATACATACTTGCAGTCACTGAGACGT TCGAGACGTCTTGGGCACTGACTGCTCCAGCAGTACATTTC). Double-stranded oligonucleotides (ds oligos) were generated by annealing the above two strands with the pcDNA™ 6.2-GW/± EmGFP-miR vector and transforming the ligated construct into competent Escherichia coli, using a BLOCK-iT pol II miRNAi expression vector and an EmGFP kit from Invitrogen (Shanghai, China). A pre-miRNA expression cassette was transferred to a Gateway® adapted destination vector utilizing Pol II promoters to form a new miRNA expression clone containing attR substrates. The vector was identified by analysing the plasmid sequence (Invitrogen, Shanghai, China). The lenti-pre-AMO-miR-1 vectors used for the experiments (2.0 μL) contained 1.0 × 10⁵ transducing units (TUs)/mL. Virus suspensions were stored at −80 °C until use and were briefly centrifuged and kept on ice immediately before injection.

Stereotactic injection of lentiviral vectors

The mice were anaesthetized with sodium pentobarbital (40 mg/kg) by intraperitoneal injection and maintained under anaesthesia using 0.5–1.0% isoflurane. The depth of anaesthesia was monitored by detection of reflexes, heart rate and respiratory rate. After anaesthesia, the mice were placed on a stereotaxic frame (RWB Life Science Co. Ltd., China). Injections were administered in the CA1 area of the hippocampus using a 5-μL Hamilton syringe with a 33-gauge tip needle at −2.2–2.5 mm below the surface of the dura. The injection coordinates relative to the bregma were as follows: anteroposterior (AP), −2.8 mm; mediolateral (ML), ±3.0 mm; and dorsoventral (DV), 2.2–2.5 (Hamilton, Bonaduz, Switzerland). The needle was maintained in place for another 2 min after injection and was then withdrawn very slowly to avoid backflow of the solution. The accuracies of the injection sites were confirmed by stereotaxic injection.
of Evans blue (Sigma Chemical Co., St. Louis, Missouri, USA) directly into the hippocampus CA1 subfield [18, 29].

Administration of GW4869 into MI and Tg mice
GW4869 (N,N’Bis[4-(4,5-dihydro-1H-imidazol-2-yl)phenyl]-3,3'-p-phenylene-bis-acrylamide dihydrochloride; MW 677.5 g/mol; Cayman Chemical) was first dissolved in the solution of DMSO as the storage solution at a concentration of 8 mg/mL, which was diluted to a concentration of 0.3 mg/mL by 0.9% NaCl before administration. The method of intraperitoneal injection of GW4869 was used in this study, and the single injection dosage was 200 μL at 0.3 mg/mL for each mouse. Equal volumes of 3.75% DMSO diluted with 0.9% NaCl were injected into mice as a control group. The first injection was performed on the day before LCA surgery for myocardial infarction mice and at the age of 6 M for Tg mice. Thereafter, GW4869 administration was performed 1 times per 2 d for 15 d, and the mice were sacrificed 24 h after the final injection for the further experiments.

Blood sampling
Whole blood samples (0.5 mL per mouse) were collected from the hearts of the anaesthetized mice (sodium pentobarbital, 100 mg/kg, i.p.) via direct venous puncture. Blood was collected into a vacuum blood collection tube containing sodium citrate and was then carefully transferred into an RNase-free tube for RNA extraction [5, 17].

Primary culture of neonatal rat hippocampal and cortical neurons (NRNs)
The primary culturing of NRNs was performed according to procedures that have been previously described in detail [30]. Briefly, hippocampal and cortical regions were collected from rat pups at postnatal day 0 (P0) and placed immediately into ice-cold PBS. The samples were then dissected and incubated in the presence of 0.125% trypsin (Gibco, USA) for 15 min in a water bath at 37 °C. After dispersion, cells were plated onto cell culture plates pre-coated with 10 μg/mL poly-D-lysine (Sigma, USA). The cells were maintained in culture media containing neurobasal medium (Gibco, USA) with 2% B27 supplement (Invitrogen, USA) and 10% foetal bovine serum (FBS, HyClone, Logan, UT) at a density of 1 × 3 × 10^5 cells/cm². The cultures were incubated in a 37 °C humidified chamber that was maintained at 5% CO₂. The cells were fed by exchanging 50% of the culture media twice a week. The neurons were treated with 5 μM cytosine arabinoside (Sigma, USA) after 3 d in culture to inhibit astrocyte proliferation. For all experiments, neurons were used at 7–10 days after plating.

Transfection procedures
A dose of 75 pmol/mL miR-1, mis-miR-1, AMO-1, Snap25-ODN siRNA or diethyl phosphorocyanidate (DEPC) water was transfected into NRNs using X-treme GENE siRNA transfection reagent (Cat.# 04476093001, Roche, USA) according to the manufacturer’s instructions [29]. The cells were collected for total RNA isolation or protein purification at 48 h post-transfection.

Dual luciferase reporter assay
We cloned the full-length of 3’UTR of the Snap25 gene to generate reporter vectors with miRNA binding sites. The full-length 3’UTR of Snap25 was then amplified by PCR and cloned into the psi-CHECK™-2 luciferase expression vector containing NotI-XhoI sites. HEK293T cells minimally expressing endogenous miR-1 were used in the luciferase assay. These cells were cultured in DMEM with 10% FBS and 100 μg/mL penicillin/streptomycin. In vitro transfection and luciferase assays were conducted using HEK293T cells (plated at 40%~50% confluence and 20 μmol/L miR-1, AMO-1, or mis-miR-1 as well as 0.5 μg psi-CHECK™-2-target DNA firefly luciferase vector, 1 μL blank plasmid and Lipofectamine 2000 (Invitrogen) transfection reagent according to the manufacturer’s instructions. After 48 h, the luciferase activity was measured with a dual luciferase reporter assay kit (Cat.#. E1910, Promega) and a luminometer (GloMax™ 20/20, Promega, USA). Nucleotide-substitution mutations in the 3’UTRs of Snap25 were generated by direct oligomer synthesis. The sequences of constructs were verified [26, 29].

TaqMan qRT-PCR
The levels of miR-1 were quantified using a TaqMan® MicroRNA Reverse Transcription Kit (Cat.#4,366,596, Applied Biosystems) and TaqMan® Gene Expression Master Mix (the target sequences included UGGAAGUAAAAGAAGUGUGUAU (Cat.#002064) for rats and UGGAAUGUAAGAAGUAAUGUAU (Cat.#002222) for mice; Applied Biosystems), and U6 (Cat.# 001973, Applied Biosystems) was used as an internal control. The TaqMan qRT–PCR probes and primers for miR-1 were designed by Invitrogen (USA). qRT-PCR was performed with a thermocycler (ABI Prism® 7500 fast, Applied Biosystems, Foster City, CA) for 40 cycles. The threshold cycle (Ct) was defined as the fractional cycle number at which the fluorescence passed the fixed threshold. PCR was performed as follows: (1) 95 °C for 10 min; (2) 95 °C for 15 s, followed by 60 °C for 1 min (repeat (2) for 40 cycles). The results were normalized against U6 expression levels using the ΔΔCt method [26].

Western blot analysis
Total protein samples for western blot analysis were extracted from cultured NRNs or the left ventricle and hippocampi of the mice after they were anaesthetized.
with sodium pentobarbital (100 mg/kg, i.p.). Mouse death was then confirmed by exsanguination according to a previously described method [31]. Hippocampi for primary cell culture were collected from neonatal Sprague-Dawley (SD) rats after the administration of 20% isoflurane and confirmation of death by cervical dislocation. Anti-SNAP-25 (1:1000, ab5666, Abcam, MA, USA), anti-VAMP-2 (1:10000, 104,211, Synaptic Systems, Gottingen, Germany) and anti-Syntaxin-1A (1:5000, 104,211, Synaptic Systems, Gottingen, Germany), anti-Munc-18 (1:1000, 116,002, Synaptic Systems, Gottingen, Germany), anti-CD63 (1:1000, Ab193349, Abcam, MA, USA), were used as primary antibodies. β-actin (1:1000, G8795, Sigma, Saint Louis, MO, USA) was selected as an internal control. The blots were detected with an Odyssey Infrared Imaging System (LI-COR Biosciences, Lincoln, NE, USA). The final results were expressed as fold changes compared with the control values.

Detection of synaptic vesicle exocytosis with FM1–43 dyes

FM1–43 fluorescence dye (T35356, Invitrogen, Oregon, USA) was dissolved in deionized, distilled water to make stock solutions (1 mg/mL) and stored at 2–8 °C. After the NRNs were cultured for 10–14 days in the cover glasses, the cell slides were placed in glass bottom of cell culture dishes (Nest, Cat. NO. 801002, China) and washed twice gently with 0.9% NaCl at 37 °C. First, the NRNs were incubated with an FM1–43 fluorescence working solution (10 μg/mL) dissolved in 0.9% NaCl for 3 min at room temperature to allow FM 1–43 binding to the outer membrane of NRNs. Then, 70 mM KCl dissolved in 0.9% NaCl was added to the FM1–43 loaded NRNs for 3 min at room temperature to internalize FM1–43 dye by endocytosis. Subsequently, the NRNs were washed again to remove the extracellular FM1–43 dye with 0.9% NaCl at 37 °C and then placed on a confocal microscope stage. The FM1–43 loaded NRNs were treated with 70 mM KCl solution to elicit synaptic vesicle exocytosis. The declined FM1–43 fluorescence dye signal due to the activated synaptic vesicle exocytosis was monitored by confocal microscopy (Olympus FV1000, Japan), and fluorescence images were acquired every 500 ms [8, 32].

Statistical analysis

The data are described as the mean ± SEM. FM1–43 staining was performed using factorial ANOVA (split-plot design) after performing Mauchly’s test of sphericity with a P > 0.05. Post hoc analyses of significant main effects were further performed using Fisher’s least significant difference test (PLSD) for multiple comparisons. The independent sample test was calculated using the Levene test for equality of variance. If P > 0.05, then an independent Student’s t-test was used for the comparison between two groups. If P < 0.05, then the Kruskal-Wallis rank sum test was performed. One way ANOVA was performed for the comparison among multivariate groups and post hoc analyses of significant main effect were further examined using Fisher’s PLSD for multiple comparisons. P < 0.05 was considered statistically significant. SAS 9.1 software (serial number: 989155; Institute Inc. China) was used for all statistical analyses.

Results

Abnormal synaptic vesicle distribution in the hippocampi of cardiac-specific miR-1 overexpression Tg mice

To observe whether the overexpression of miR-1 in the heart could induce synaptic pathological remodelling in the brain, a mouse line for the cardiac-specific overexpression of miR-1-2 driven by the α-myosin heavy chain (α-MHC) promoter was developed as described previously [26]. Similar to the previous study [18], miR-1 expression was significantly increased in both the hearts (Fig. 2a) and hippocampi (Fig. 2b) of the Tg mice compared with that in age-matched wild type (WT) mice, as validated by in situ hybridization technology. Using transmission electron microscopy (TEM) examination, we found that the total number of synaptic vesicles (SVs) in the whole presynaptic area of the hippocampus was significantly increased (Fig. 2c and d). Interestingly, we found that SVs at 0 nm distances from the active zone were the same in the hippocampi of the mice in the two groups but they were increased at 50 and 100 nm distances far from the active zone (Fig. 2c and e). However, no difference was observed in the active zone length between the groups (Fig. 2f). These results suggest a blunted fusion process between synaptic vesicles and presynaptic membranes in the hippocampi of the Tg mice. As displayed in Fig. 2g, SV exocytosis is a complex process that is dependent on the cooperation of multiple proteins including the SNARE complex, which is assembled by SNAP-25 (synaptosomal-associated protein 25), VAMP-2 (vesicle associated membrane protein 2, also known as synaptobrevin-2) and syntaxin-1 (also known as HPC-1), which determines the presynaptic release probability by controlling the fusion kinetics [7, 32]. We hence evaluated the expression of SNARE-complex proteins. We found that the SNAP-25 and syntaxin-1A protein levels were significantly reduced, while VAMP-2 expression was unchanged (Fig. 2h & i). Furthermore, the protein level of Munc-18, which is considered to play a key role in presynaptic vesicle exocytosis by interacting with the SNARE complex [7], was unchanged in the hippocampi of the Tg mice compared with WT mice, (Fig. 2h & i).

MIR-1 targets the Snap25 gene and attenuates synaptic vesicle exocytosis

We next aimed to clarify whether SNAP-25 and syntaxin-1A are the targets of miR-1. By performing computational analysis of miRNA databases (TargetScan...
5.1 and miRanda), we found that there is a putative miR-1 binding site in the 3’UTR of the Snap25 gene at the site between 413 bp and 420 bp (Fig. 3a). However, we did not find the binding site of miR-1 on the Stx1a gene (encoding Syntaxin-1A protein). To experimentally verify this finding, we first cloned the full-length 3’UTR of Snap25 into a luciferase-expressing reporter vector (Fig. 3b) and then evaluated the effects of miR-1 on reporter activity in HEK293T cells. The data showed that the co-transfection of the psiCHECK™-2 vector plasmid with miR-1 consistently resulted in a reduction of luciferase activity compared with that observed in transfection of the plasmid alone (Fig. 3c, Renilla/firefly: 1.42 ± 0.08 in blank vs. 0.45 ± 0.03 in miR-1, P < 0.05). Application of a 2′-O-methyl antisense oligoribonucleotide to miR-1 (AMO-1), a specific inhibitor of miR-1, eliminated the repressive effect of miR-1 on the Renilla fluorescence signal (Fig. 3c, Renilla/firefly: 1.52 ± 0.116 in AMO-1 vs. 0.45 ± 0.03 in miR-1, P < 0.05). However, the negative control of both miR-1 (mis-miR-1) and AMO-1 (mis-AMO-1) failed to influence miR-1 action on the Renilla fluorescence signal (Fig. 3c). Furthermore, mutating the binding sites of the Snap25 gene abolished the effect of miR-1 (Fig. 3d), suggesting that these binding sites contribute to the repressive effects of miR-1.

Fig. 2 Abnormal synaptic vesicle distribution and related protein expression in the hippocampi of cardiac-specific miR-1 overexpression Tg mice. a-b Elevation of miR-1 levels in the hearts (a) and hippocampi (b) of Tg mice compared with WT mice by in situ hybridization (ISH) using DIG-labelled mmu-miR-1 probes and scramble-miR-1 (mis-miR-1) probes. The positive signal for mmu-miR-1 probes was stained in blue. c Representative electron micrographs of Tg mice. d Morphometric analysis of the total number of SVs in the synapses of neurons from the hippocampi of Tg and WT mice. P<0.0001, Kruskal-Wallis rank sum test: P < 0.0001. n = 20 synapses from 3 mice for each group. e Quantitative analysis of the distribution of SVs in WT and Tg nerve terminals. 0 nm shell: P<0.0001; 50 nm shell: P = 0.001; 100 nm shell: P<0.0001, Kruskal-Wallis rank sum test: P < 0.0001. n = 20 synapses from 3 mice for each group. f Active zone length was not different between WT and Tg mice. P=0.016, Kruskal-Wallis rank sum test: P = 0.844. n = 20 active zones from 3 mice for each group. g Schematic diagram of synaptic vesicle fusion. Under the stimulation of Ca²⁺, Munc-18 switches and binds to syntaxin-1 and assembles SNARE proteins into SNARE-complexes that promote SV fusion with presynaptic membranes. Syntaxobrevin 2: VAMP-2. h Representative immunoblotting images of VAMP-2, SNAP-25, Syntaxin-1A, Munc-18. i Statistical analysis of VAMP-2: P=0.478, Student’s t test: t = 0.200, P = 0.845; SNAP-25: P=0.562, Student’s t test: t = 3.463, P = 0.006; Syntaxin-1A: P=0.670, Student’s t test: t = 5.067, P < 0.0001. n = 6; Munc-18: P=0.706, Student’s t test: t = −0.656, P = 0.527. n = 6. *P < 0.05; SV: synaptic vesicle
To observe whether miR-1 regulates the expression of SNAP-25, we transfected miR-1 and/or AMO-1 directly into NRNs using the X-treme GENE siRNA transfection reagent. The successful transfection of miR-1 and/or AMO-1 was confirmed by miR-1 levels in NRNs detected by qRT-PCR (Fig. 4a). Based on this condition, by immunoblotting analysis, we found that miR-1 effectively inhibited SNAP-25 protein expression by nearly 30% relative to the level of the control group, whereas the scrambled negative control mis-miR-1 failed to affect the protein level (Fig. 4b). In contrast, AMO-1 rescued the miR-1-mediated downregulation of SNAP-25 (Fig. 4b).

As predicted, the overexpression of miR-1 did not regulate syntaxin-1A expression in NRNs after transfection with miR-1 and/or AMO-1 (Fig. 4c).

To observe whether miR-1 could influence SV number and distribution in vitro, TEM detection was performed. We observed a significantly increased number of SVs in the active zone of NRNs after the transfection of miR-1 for 48 h, however, co-transfection of AMO-1 with miR-1 blocked the miR-1-mediated aggregation of synaptic vesicles inside the synapses (Fig. 4d).

FM1-43 fluorescence dyes are widely used to image vesicle exocytosis and endocytosis [8]. We observed that the FM1-43 fluorescent signal in NRNs transfected with mis-miR-1 gradually decreased after 70 mM KCl stimulation (Fig. 4e). The ratio of F/F0 was reduced to 0.5 at 5 s, and further decreased to 0.4 at 10 s after KCl stimulation (Fig. 4f). The transfection of
miR-1 markedly blocked the attenuated FM1–43 fluorescent signal induced by KCl, which was reversed by the co-transfection of AMO-1 (Fig. 4e & f).

To clarify whether the observed changes in SNAP-25 expression and exocytosis in cultured NRNs are the direct results of miR-1 action on the binding sites of this
genes, an miRNA-masking antisense ODN fragment (miR-mask) was designed from the 3’UTR of the Snap25 gene sequence. The miR-mask was named Snap25-ODN, which acts on a specific binding site between 408 ~ 430 bp of this gene and minimally influences the effects of miR-1 on other target genes and other binding sites of this gene [33]. As expected from their principle action, Snap25-ODN, unlike AMO-1, did not affect the miR-1 level in cells co-transfected with miR-1 (Fig. 5a). As predicted, Snap25-ODN inhibited the action of miR-1 on SNAP-25 expression (Fig. 5b), suggesting that miR-1 post-transcriptionally regulated SNAP-25 expression by binding to a site located at 413 ~ 420 bp in the 3’UTR of the Snap25 gene. Furthermore, Snap25-ODN successfully prevented the miR-1 overexpression-induced impairment of vesicle exocytosis (Fig. 5c & d). All these results suggested that miR-1 overexpression triggered impaired synaptic vesicle exocytosis, mainly by reducing the SNAP-25 protein level.

Furthermore, to determine whether miR-1 is involved in the declined synaptic vesicle exocytosis in the Tg mice, anti-miR-1 oligonucleotide fragments carried by a lentivirus vector (lenti-pre-AMO-1) were stereotaxically injected directly into the CA1 area of the hippocampus bilaterally in the Tg mice at 5 months. The miR-1 levels in the hippocampi of the Tg mice were successfully minimized...
inhibited by lenti-pre-AMO-1 after injection for 2 months (Fig. 6a). Similar to observations in the in vitro experiments (Fig. 4b), a reduction in SNAP-25 protein levels was observed in the hippocampi of the Tg mice with lenti-pre-mis-AMO-1 which was effectively prevented by lenti-pre-AMO-1 injection (Fig. 6b). Furthermore, lenti-pre-AMO-1 significantly reversed the increased SV number in the Tg mice (Fig. 6c & d) and attenuated SV redistribution at the presynaptic active zone in the hippocampi of the Tg mice (Fig. 6c & e). In this way, lenti-pre-AMO-1 injection did not affect the active zone length compared with that in the uninjected Tg mice (Fig. 2f). These data indicated that the hippocampal overexpression of \( \text{miR-1} \) impaired SV exocytosis in Tg mice was associated with the decreased expression of SNAP-25 protein in the hippocampi.

The cardiac \( \text{miR-1} \) overexpression-mediated attenuation of SV exocytosis involves exosomes

We next examined why the overexpression of \( \text{miR-1} \) in the heart affects SV exocytosis in the brain. Studies have reported that miRNAs could be secreted into the extracellular space of donor cells and then transported and accepted by recipient cells with functional targeting capabilities [3, 11]. As one kind of miRNA transporter, exosomes not only contain up to 121 miRNAs, including \( \text{miR-1}, \text{miR-15}, \text{miR-16}, \text{miR-17}, \text{miR-18}, \text{miR-181}, \text{miR-375}, \text{lin-4}\) and \( \text{let-7} \) [14], but also show an increase in the serum of the infarcted heart of rats [19, 34]. We hence speculated that exosomes might participate in this process as transporters. To elucidate this issue, we applied GW4869 by intraperitoneal injection to inhibit sphingomyelinase, which was reported to inhibit

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**Fig. 6** Stereotaxic injection of lenti-pre-AMO-1 into the hippocampi of Tg mice improved SV distribution. **a** Stereotaxic injection of lenti-pre-AMO-1 into the CA1 domain reduced the miR-1 level in the hippocampi of Tg mice as determined by qRT-PCR analysis. \( P_{\text{Levene}} = 0.01 \), one-way ANOVA: \( F = 10.872, P = 0.01 \); Fisher's PLSD test: \( P_{\text{WT:Tg}} = 0.005, P_{\text{Tg+lenti-pre-mis-AMO-1:Tg+lenti-pre-AMO-1}} = 0.012 \). \( n = 3 \) batches of cells for each group. **b** Stereotaxic injection of lenti-pre-AMO-1 into the hippocampi of Tg mice derepressed the expression of SNAP-25 protein. \( P_{\text{Levene}} = 0.004 \), one-way ANOVA: \( F = 15.700, P < 0.001 \); Fisher's PLSD test: \( P_{\text{WT:Tg}} = 0.001, P_{\text{Tg+lenti-pre-mis-AMO-1:Tg+lenti-pre-AMO-1}} < 0.0001 \). \( n = 6 \) animals for each group. **c** Representative electron micrographs showing the effect of stereotaxic injection of lenti-pre-AMO-1 on SVs distribution at presynaptic active zone of Tg mice. **d** Lenti-pre-AMO-1 prevented an increase in the total number of SVs in the hippocampal synapses of Tg mice. \( P_{\text{Levene}} < 0.003 \), one-way ANOVA: \( F = 33.161, P < 0.0001 \); Fisher's PLSD test: \( P_{\text{WT:Tg}} < 0.0001, P_{\text{Tg+lenti-pre-mis-AMO-1:Tg+lenti-pre-AMO-1}} < 0.0001 \). \( n = 20 \) active zones from 3 mice for each group. **e** Lenti-pre-AMO-1 redistributed SV location in Tg mice nerve terminals. 0 nm shell: \( P_{\text{Levene}} < 0.001 \), one-way ANOVA: \( F = 2.928, P = 0.062 \); Fisher's PLSD test: \( P_{\text{WT:Tg}} = 0.596, P_{\text{Tg+lenti-pre-mis-AMO-1:Tg+lenti-pre-AMO-1}} = 0.024 \). 50 nm shell: \( P_{\text{Levene}} = 0.044 \), one-way ANOVA: \( F = 7.517, P = 0.01 \); Fisher's PLSD test: \( P_{\text{WT:Tg}} = 0.003, P_{\text{Tg+lenti-pre-mis-AMO-1:Tg+lenti-pre-AMO-1}} = 0.001 \). 100 nm shell: \( P_{\text{Levene}} = 0.005 \), one-way ANOVA: \( F = 11.161, P < 0.0001 \); Fisher's PLSD test: \( P_{\text{WT:Tg}} = 0.014, P_{\text{Tg+lenti-pre-mis-AMO-1:Tg+lenti-pre-AMO-1}} < 0.0001 \). \( n = 20 \) synapses from 3 mice for each group. **f** Lenti-pre-AMO-1 did not affect the length of active zone. \( P_{\text{Levene}} = 0.551 \), one-way ANOVA: \( F = 0.404, P = 0.670 \); Fisher's PLSD test: \( P_{\text{WT:Tg}} = 0.940, P_{\text{Tg+lenti-pre-mis-AMO-1:Tg+lenti-pre-AMO-1}} = 0.419 \). \( n = 20 \) active zones from 3 mice for each group. *\( P < 0.05 \) versus WT; #\( P < 0.05 \) versus Tg.
exosome generation [17]. As illustrated in Fig. 7a, we observed that GW4869 significantly inhibited the reduction in CD63, a marker of exosomes, in Tg mouse hearts, suggesting that the generation of exosomes in the heart was prevented. As predicted, the miR-1 level in the blood (Fig. 7b) and hippocampi (Fig. 7c) of Tg mice were both decreased in Tg mice after GW4869 intraperitoneal injection. Notably, the injection of GW4869 significantly reversed the reduction in SNAP-25 protein in the hippocampi of Tg mice (Fig. 7d).

Importantly, miR-1 has been reported to be increased in the blood of both acute myocardial infarction (AMI) rats and patients [5, 35, 36] and hippocampi of myocardial infarction (MI) mice [19]. Could MI also induce impaired presynaptic plasticity and share the same mechanism as that in Tg mice? We first developed an MI animal model by ligating of the left coronary artery (LCA) for 30 days. Similar to the observation in experiments of Tg mice (Figs. 2c & 6c), the number of SVs was significantly increased in MI mice compared with that in sham mice (Fig. 8a & b), and SVs were more concentrated at sites of 50 nm and 100 nm shells from the active zone in MI mice compared with those in age-matched sham mice (Fig. 8a and c). Strikingly, all these changes were prevented by intraperitoneal injection of GW4869 (Fig. 8a-c). However, neither MI nor GW4869 affected the active zone length (Fig. 8d). We then assessed the expression of synaptic fusion-related protein in the hippocampi of MI mice. We found that, consistent with Tg mice, both Munc-18 and VAMP-2 protein levels were unchanged compared with those in the sham control (Fig. 8e). However, SNAP-25 protein expression was significantly reduced from the 15th to the 30th day after LCA (Fig. 8e). Interestingly, although

![Fig. 7](image)

**Fig. 7** Inhibition of exosome generation derepressed the expression of SNAP-25 protein in the hippocampi of Tg mice. **a** Intraperitoneal injection of GW4869 inhibited CD63 expression in the heart of Tg mice at the age of 6 M compared with age-matched Tg mice, as measured by western blotting. $P_{\text{Levene}} = 0.419$, one-way ANOVA: $F = 12.778$, $P = 0.001$; Fisher's PLSD test: $P_{\text{WT:Tg}} = 0.002$, $P_{\text{Tg::GW4869}} = 0.001$. $n = 5$ animals for each group. **b** Intraperitoneal injection of GW4869 prevented the elevation of miR-1 level in the blood (b: $P_{\text{Levene}} < 0.001$, one-way ANOVA: $F = 26.140$, $P < 0.0001$; Fisher's PLSD test: $P_{\text{WT:Tg}} < 0.0002$, $P_{\text{Tg::GW4869}} < 0.0001$. $n = 6$ animals for each group). **c** Intraperitoneal injection of GW4869 prevented the elevation of miR-1 level in the hippocampi (c: $P_{\text{Levene}} < 0.001$, one-way ANOVA: $F = 7.756$, $P = 0.011$; Fisher's PLSD test: $P_{\text{WT:Tg}} = 0.004$, $P_{\text{Tg::GW4869}} = 0.0181$. $n = 4$ animals for each group) of Tg mice, as measured by qRT-PCR. **d** Intraperitoneal injection of GW4869 derepressed SNAP-25 expression in the hippocampi of Tg mice, as measured by western blotting. $P_{\text{Levene}} = 0.419$, one-way ANOVA: $F = 12.778$, $P = 0.001$; Fisher's PLSD test: $P_{\text{WT:Tg}} = 0.002$, $P_{\text{Tg::GW4869}} = 0.001$. $n = 6$ animals for each group. *$P < 0.05$ versus WT; # $P < 0.05$ versus Tg.
Fig. 8 Inhibition of exosome generation improves MI disrupted SV distribution. a Intraperitoneal injection of GW4869 prevented the synaptic vesicles accumulation at the presynaptic active zone in hippocampal neurons of MI mice. b GW4869 prevented an increase of the total SV number in the synapses in the hippocampi of MI mice. P_{Levene} < 0.0001, one-way ANOVA: F = 19.799, P < 0.0001; Fisher’s PLSD test: P_{sham, MI} < 0.0001, P_{GW4869, MI} < 0.0001. n = 20 synapses from 3 mice for each group. c GW4869 redistributed SVs at the active zones of synapses in the nerve terminals of Tg mice. 0 nm shell: P_{Levene} = 0.669, one-way ANOVA: F = 3.424, P = 0.039; Fisher’s PLSD test: P_{sham, MI} = 0.360, P_{GW4869, MI} = 0.012; 50 nm shell: P_{Levene} = 0.016, one-way ANOVA: F = 6.703, P = 0.002; Fisher’s PLSD test: P_{sham, MI} = 0.003, P_{GW4869, MI} = 0.002; 100 nm shell: P_{Levene} < 0.0001, One way ANOVA: F = 14.785, P < 0.0001; Fisher’s PLSD test: P_{sham, MI} = 0.001, P_{GW4869, MI} < 0.0001. n = 20 synapses from 3 mice for each group. d GW4869 did not affect the length of active zone in hippocampi of MI mice. P_{Levene} = 0.271, one-way ANOVA: F = 1.084, P = 0.345; Fisher’s PLSD test: P_{sham, MI} = 0.185, P_{GW4869, MI} = 0.885. n = 20 active zones from 3 mice for each group. e Expression of SV related proteins in the hippocampi of MI mice. Left: sample bands of immunoblotting images. Right: statistical analysis of Munc18, VAMP-2, syntaxin-1A and SNAP-25. *P < 0.05 (f) Intraperitoneal injection of GW4869 prevented the elevation of miR-1 level in the hippocampi of MI mice, as measured by qRT-PCR. P_{Levene} < 0.0001, one-way ANOVA: F = 7.756, P = 0.011; Fisher’s PLSD test: P_{sham, MI} = 0.004, P_{GW4869, MI} = 0.018. n = 4 animals for each group. g Intraperitoneal injection of GW4869 prevented the decreased SNAP-25 expression in the hippocampi of MI mice, as measured by western blotting. P_{Levene} = 0.265, one-way ANOVA: F = 7.647, P = 0.011; Fisher’s PLSD test: P_{sham, MI} = 0.010, P_{GW4869, MI} = 0.005. n = 4 animals for each group. *P < 0.05 versus WT, #P < 0.05 versus Tg.
the syntaxin-1A protein level was decreased in the hippocampi of MI mice on the 30th day of LCA, the level was not changed before 15 d (Fig. 8e). Importantly, the intraperitoneal injection of GW4869 into MI mice markedly prevented the increased miR-1 levels (Fig. 8f) and decreased SNAP-25 expression (Fig. 8g) in the hippocampi of MI mice. All these data suggested that the cardiac miR-1 overexpression-mediated attenuation of synaptic vesicle exocytosis involves exosomes.

Discussion
In the present study, we reported that miR-1 overexpression in the heart could result in the attenuation of SV exocytosis, which is controlled by increased miR-1 levels in the hippocampus originating from exosome-mediated transportation from the heart to the brain through circulation. The molecular mechanism involves miR-1 targeting the 3'UTR of the Snap25 gene and post-transcriptionally inhibiting the expression of SNAP-25 protein (Fig. 9). This study improved our understanding of how heart disease induces brain dysfunction at the miRNA level.

The link between cardiac diseases and cognitive deterioration has been accepted from the concept of “cardiogenic dementia” which was proposed in the late 1970s [37]. However, most studies have focused on cardiac disease-induced chronic brain hypoperfusion (CBH) due to the reduction of cardiac output [22, 24, 38]. Whether heart abnormalities alone could lead to brain damage and even affect cognition is elusive. Using the cardiac-specific overexpression of miR-1-2 in transgenic mice driven by α-MHC, our previous study demonstrated that the overexpression of miR-1 in the hearts of Tg and MI mice could increase its expression in the hippocampus and impair cognition by post-transcriptionally inhibiting BDNF expression [18, 39].
Because synaptic pathology is a prominent feature of Alzheimer's disease (AD) or vascular dementia (VaD) [20], we propose that the cardiac overexpression of miR-1 affects synaptic plasticity. Using TEM technique, we observed an increase in miR-1 levels and SV redistribution in the hippocampi of Tg mice, and this phenomenon was reversed by the stereotaxic injection of AMO-1. This result suggests that the cardiac-specific overexpression of miR-1 could impair synaptic plasticity, which may associate with increased miR-1 levels in the hippocampus. To verify this hypothesis, we performed in vitro experiments and demonstrated that the overexpression of miR-1 led to more synaptic vesicle retention at presynaptic synaptosomes and attenuated KCl-induced FM1–43 fluorescent signal decline, which were prevented by the miR-1 antagonist AMO-1. The decisive step of presynaptic vesicle exocytosis is Ca^{2+}-triggered fusion between synaptic vesicles and presynaptic membranes, which is mediated by the interaction of SNAREs and SM proteins containing SNAP-25, VAMP-2, syntaxin-1A and Munc-18 [7]. Here, we found that both SNAP-25 and syntaxin-1A protein expression levels were decreased. To clarify the molecular mechanism, we used three strategies, including the screening of the bioinformatics database, and in vitro antisense antagonist and miR-masking techniques, and found that miR-1 directly regulates SNAP-25 expression but has no regulatory effect on syntaxin-1A protein expression. Interestingly, a previous study using SNAP-25-deficient neuron reported that the knockdown of SNAP-25 does not modify fusion pore opening or the rate of endocytosis to a degree that could alter Ca^{2+}-triggered release and FM dye release kinetics significantly [40]. In the present study, we observed that miR-1 gain of function could induce a decline in synaptic vesicle exocytosis, which was prevented by shielding the function of SNAP-25 via masking the miR-1 binding site in the 3’UTR of the Snap25 gene located at the site of 413 ~ 420 bp. All these studies implied that miR-1 could induce the impairment of synaptic vesicle exocytosis by inhibiting the expression of SNAP-25 protein. In spite of the fact that SNAP-25 is part of the basic core fusion machinery, the role of SNAP-25 in excitatory- or inhibitory- neurotransmission is controversial [40–43]. Although we were not able to assess how miR-1 or SNAP-25 affects glutamate release or aminobutyric acid neurotransmission, we believe that it is possible that miR-1 or SNAP-25 play a role in both excitatory and inhibitory neurotransmission. However, this hypothesis needs to be studied further. In addition, vesicle release is a cascade response process involving a battery of proteins including synapsins, synaptotagmin-1, Munc-13, Munc-18, complexins and SNAREs proteins. In this study, we focused on SNARE proteins. Whether the other vesicle-related proteins are also involved in MI-mediated presynaptic plasticity needs to be studied further.

Our previous study demonstrated that the increased miR-1 in the hippocampi of cardiac-specific miR-1-overexpressing Tg mice was not due to endogenous biogenesis, but was associated with exosomes that mediated miR-1 transport from the heart to the brain [18, 19]. Importantly, exosomal miR-1 was increased in both the heart and the blood [34, 44]. Because the physiological process of exosomes as transporters is to release the transported substances into recipient cells by fusing with the recipient cell membrane and then becoming part of the cell membrane, the exosomal miR-1 in the hippocampus might not represent the transported miR-1 level from the heart. In this study, to evaluate the role of cardiac derived exosomes in regulating hippocampal miR-1 levels, we performed an intraperitoneal injection of GW4869 to prevent exosome biogenesis in the heart. We found that GW4869 not only inhibited the expression of CD63 in the heart, a marker of exosomes [17], but also inhibited the increased miR-1 level in both the blood and hippocampi of Tg mice and improved the decreased SNAP-25 expression and SV distribution at the active zone in the hippocampi of Tg mice. In addition, if miR-1 can travel through blood exosomes, then this molecule may reach other brain areas, such as the frontal cortex and even the whole brain, most likely downregulating SNAP-25. This study focused on the hippocampus because the hippocampus is the main brain area associated with spatial memory, especially at the early stage of AD. MI is also considered a risk factor for the onset of AD and may be planned into the preclinical stage of AD. Our previous study demonstrated that miR-1 Tg mice displayed an impairment of spatial memory. However, whether the overexpression of miR-1 in the heart could also induce morphological or functional remodelling in other brain areas and what is the molecular mechanism is very interesting and needs to be clarified further.

**Conclusions**

Taken together, our findings demonstrate that the overexpression of miR-1 in the heart attenuated synaptic vesicle exocytosis in the hippocampus by posttranscriptionally regulating SNAP-25 through exosome transportation (Fig. 9). This study improved our understanding of the relationship between cardiovascular disease and brain dysfunction at the miRNA levels.

**Abbreviations**

AMO-1: anti-miR-1 oligonucleotide fragment; CBH: chronic brain hypoperfusion; miRNAs: microRNAs; NRNs: neonatal rat hippocampal and cortical neurons; ODN: oligodeoxynucleotides; SNAP-25: synaptosome-associated protein 25; SNAP: soluble N-ethylmaleimide-sensitive factor attachment protein receptor; Tg: transgenic
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Availability of data and materials

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

Authors’ contributions

JA designed, supervised and involved in drafting and editing the manuscript. MJD, MLY, QW, MM, DS, LLS, KXL, YQ, QS, XYZ, SUH, JCM and TB performed the experiments. JA and TB performed statistical analyses. All authors read and approved the final manuscript.

Consent for publication

Not applicable

Competing interests

The authors declare that they have no competing interests.

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