Zinc finger E-Box binding homeobox 2 (ZEB2)-induced astrogliosis protected neuron from pyroptosis in cerebral ischemia and reperfusion injury

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Introduction

Stroke, including ischemic stroke and hemorrhagic stroke, is currently one of the leading causes of human disability and mortality worldwide, of which 87% are ischemic stroke caused by obstruction of the cerebral arteries [1,2]. When ischemic stroke occurs, impaired blood supply to the brain tissue, leading to ischemia and hypoxia, which in turn leads to clinical manifestations of the corresponding neurological deficits, thus resulting in disability, or even death of the patient after an ischemia stroke [3,4]. Recently, to release vascular occlusion and restore cerebral perfusion, intravenous thrombolysis and mechanical thrombectomy are the two main strategies to achieve cerebral blood flow recanalization [5]. However, both the clinical applications are restricted by a narrow therapeutic window (only 4.5–6 h of intravenous thrombolysis and no more than 24 h of mechanical thrombectomy). Therapy beyond the therapeutic window can create reperfusion injury and aggravate the degree of brain tissue damage [6]. Neuroprotective therapies, as an adjunctive therapies to revascularization therapy, can increase the survival of nerve cells and promote the recovery of nerve function [7]. However, there has still been no successful clinical translation of neuroprotective agents to promote neuronal survival in stroke patients [8,9]. Thus, it is urgent to uncover the mechanisms of neuroprotection and neuron regeneration which can be useful in the development of new treatment methods.

In the conventional knowledge, apoptosis accounts for a significant proportion of neuronal death after cerebral ischemia [10]. Recently, however, studies found that various cell death
mechanisms, including pyroptosis, also play roles in cerebral ischemia injury [11,12]. Pyroptosis is a pro-inflammatory programmed cell death that mainly mediated by inflamma-
some of nucleotide oligomerization domain like receptors (NLRs) and depends on Caspase family [13]. It is manifested by cellular disten-
sion until the cell membrane ruptures, leading to the release of cellular contents and thus activating a strong inflammatory response [14]. According to the activation mechanism, pyroptosis can divided into caspase-1 depend-
ent and non-caspase dependent pathway [15]. Both pathways are active by cleavage of Gasdermin-D (GSDMD), an effector of pyro-
ptosis, and therefore allows the release of N-terminal pore-forming domain, which insert into the plasma membrane, forming pore that induce plasma membrane rupture and release of intracellular contents including interleukin (IL) – 1β and IL – 18 [16]. Poh et al. provided the evidence that the NLR-family CARD-
containing protein 4 (NLRC4) inflammasome complex mediates the apoptosis and pyroptosis of microglial cells under ischemic conditions [17]. Liang et al. confirmed that inhibition of Triggering Receptor Expressed on Myeloid Cells 1 (TREM-1) protects against ischemia-
induced neuronal damage and alleviates micro-
glial mediated neuro-inflammation by reducing oxidative stress and pyroptosis which observed in neuron, astrocytes, and microglia after cere-
bral ischemia conditions [18]. However, little is known the precise role of pyroptosis in cerebral injury under ischemic conditions.

Astrocytes are an important component of the central nervous systems (CNS), serving as support and compartmentalization of nerve cells and partic-
ipating in the formation of the blood-brain barrier [19]. Unlike the peripheral nervous system, axons fail to regenerate effectively after CNS injury in adult mammals, often resulting in irreversible loss of function [20,21]. Astrocytes play important regu-
larly roles in axonal regeneration during the pathophysiology of CNS injury [21]. When CNS injury occurs, it evokes a process of change in astrocyte, which is defined as astrogliosis, including processes of hypertrophy and thickening [22]. In order to protects the surrounding tissue from further damage, activated astrocytes adjacent to the lesion are tightly organized to forms a dense gelatinous scar around the wound, preventing the spread of inflammatory cells or pathogens to normal tissues and maintaining the homeostasis of the internal environment [23]. In addition, astrogliosis also promote blood-brain-barrier repair, attenuate osmotic and oxidative stresses, and thus promote axon regeneration [24]. Studies confirmed that dis-
ruption of the astrocyte responded to injury increases the size of the lesion and exacerbates the recovery of neurological function [25–27]. Prevailing evidence suggested that Zinc Finger E-Box Binding Homeobox 2 (ZEB2) is a significant factor in astrogliosis, the overexpres-
sion of ZEB2 can improve neuroprotection and recovery of neurological function in spinal cord injury and cerebral ischemic stroke by evoking astrogliosis [28]. ZEB2 is a zinc-finger homeodo-
main transcription factor protein belong to ZEB family, ZEB2 play an essential role in neuronal development [29], also participate in Epithelial-
mesenchymal transition (EMT) of cancer cell [30–33], wound healing [34] and neural development in adult [29,35]. ZEB2 was found up-regulated in cer-
ebral ischemia reperfusion injury rats, further increased expression of ZEB2 attenuated the cerebral ischemia reperfusion injury, whereas knock-
down of ZEB2 increased lesion size of cerebral ischemia reperfusion [28]. However, the role of ZEB2 in neural regeneration after cerebral ischemia reperfusion is unknown.

In this study, we hypothesized that astrocyte-
derived ZEB2 enhanced the astrogliosis and hence promoted neural regeneration. Therefore, our goal is to investigate the relationship between ZEB2 expression in astrocytes and neural regeneration in vivo and in vitro. We found that ZEB2 is a regulator of astrocytes activity, which can evoke subsequent neural regeneration by suppressing pyroptosis.

Materials and methods

Primary cell culture of neuron and astrocytes and plasmids transfected

Primary hippocampal neurons and astrocytes were isolated from Sprague Dawley (SD) rat
hippocampus as described [36,37]. In brief, SD fetal rats at 18–20 days of gestation were selected as the source of primary hippocampal cell culture. The pregnant rats were anaesthetized with ether, and the fetuses were dissected and decapitated one by one. After decapitation, the whole brain tissue was stripped out in iced Hibernate-E solution and the hippocampus was stripped bilaterally under a microscope. Neurons were cultured on poly-D-lysine coated glass coverslips in NbActiv4 (BrainBits, USA) medium containing antibiotic-antimycotic at 37°C. Passage 5–7 were used for experiment. Astrocytes were cultivated in Basal Eagle medium containing 10% fetal bovine serum, 0.45% glucose, 1 mM sodium pyruvate, 2 mM glutamate supplement and antibiotic-antifungal agent. Passage 3–5 were used for experiment.

Cells were divided into 4 groups: Ctrl group (astrocytes and neurons were co-cultured under normal condition), oxygen-glucose deprivation/reoxygenation (OGD/R) group (astrocytes and neurons were co-cultured under Glucose-deficient anaerobic condition), OGD/R + negative control (NC) group (astrocytes were transfected with pcDNA3.1-NC and then co-cultured with neurons under Glucose-deficient anaerobic condition), and OGD/R + ZEB2 group (astrocytes were transfected with pcDNA3.1-ZEB2 and then co-cultured with neurons under Glucose-deficient anaerobic condition). AS for cell transfection, pcDNA3.1-ZEB2 and pcDNA3.1-NC were synthesized from Genomeditech (Shanghai, China). Astrocytes were cultured in normal dulbecco’s modified eagle medium (DMEM) medium (Sigma China) under normal condition, pcDNA3.1-ZEB2 and pcDNA3.1-NC were transfected in astrocytes, respectively. After 48 hours, astrocytes were collected and inoculated in upper chamber of transwell with glucose-free DMEM (Sigma China) and neurons were inoculated in lower chamber with glucose-free DMEM (Sigma China). And then transwell inserts were placed in an anaerobic chamber (95% N₂, 5% CO₂ and 1% O₂) at 37°C for 4 hours. Thereafter, cell culture medium was replaced into normal DMEM medium under normal condition for 48 hours for reperfusion.

**Cell viability assay**

Cell viability of astrocytes and neurons were detected by the Cell Counting Kit-8 (CCK8) (abcam, China) [38]. After OGD/R treatment, cells (1 × 10⁴ per well) were inoculated in 96-well plates with normal DMEM medium and 10 μL CCK8 solutions were added in each well for 2 hours at 37°C, then cell viability will determine by spectrometry at 450 nm after incubation.

**Cell apoptosis assay**

Cell apoptosis of astrocytes and neurons was assessed using Annexin V-FITC Apoptosis Detection Kit (Sigma China) [39], according to the instruction of manufacturer. After stimulation with OGD/R condition, cells were inoculated in 6-well plates and 100 μL of 1 × 10⁶ cells/mL cell suspension were stained with 5 μL of Annexin V-FITC and 10 μL of Propidium iodide (PI) for 15 min at 37°C in dark, then the annexin V-FITC binding were assessed the flow cytometry at an excitation wavelength of 488 nm.

**Immunofluorescence staining [28]**

Cells fixed in 4% paraformaldehyde for 15 min. After fixation, cells permeabilized with 0.3% triton X-100 for 30 min, blocked by 10% goat serum for 60 min, then incubated with primary antibodies 5-bromo-2-deoxyuridine (BrdU) (1:250, ab6326, abcam), Neuronal nuclei (NeuN) (1:1000, ab104224, abcam), glial fibrillary acidic protein (GFAP) (1:500, ab33922, abcam), ZEB2 (1:5000, bs-20,485 R, Bioss), brain-derived neurotrophic factor (BDNF) (1:500, ab108319, abcam) at 4°C overnight. After that, the cells were incubated with secondary antibody including Fluorescein at room temperature for 2 hours in dark. All nuclei were stained with 4'-6-diamidino-2-phenylindole (DAPI) for 2 minutes in dark. Image was performed on a fluorescence microscope.

**Intracerebroventricular injection**

SD rats (grand SPF, weighing 240–270 g) were purchased from the department of experimental animals, Kunming Medical University (Yunnan,
China. Adeno-associated virus (AAV) included GFAP – specific promoters were used to pack plasmids and synthesized by HanBio, China. Rats were fasted 12 hours before surgery, and anesthetized by 42 mg/kg sodium pentobarbital, and they were fixed on the stereotaxic apparatus after they completely unconscious and the hair was removed. An incision was made along with midline scalp to expose the bregma, 1 μL of AAV-ZEB2 or its negativated control vector (1 × 10⁷ TU/mL, at a rate of 0.5 μL/min) was injected into right lateral ventricle [40]. The vectors were injected 20 days prior to the established MCAO/R model. Stereotaxic coordinates were as follows: Anteroposterior, 0.8 mm; Mediolateral, 1.5 mm; Depth, 3.5 mm.

Middle cerebral artery occlusion-reperfusion (MCAO/R) model establishment

This laboratory animal ethics (KMMU2020502, 20,200,815) has been approved by the Kunming Medical University ethics committee. The experimental procedures strictly followed the guidelines for the management and the usage of laboratory animals formulated by the national institutes of health according to the previous studies [41,42].

All rats were divided into 3 groups: sham group (n = 6), MCAO/R group (n = 6) and MCAO/R + ZEB2 group (n = 6). In brief, rats were anesthetized by injection of sodium pentobarbital (42 mg/kg), and they were fixed on the stereotaxic apparatus after they have become completely unconscious. After hair removed, an incision was made along with midline cervical to exposed the right common carotid artery (CCA), internal carotid artery (ICA) and external carotid artery (ECA). CCA and ECA were tied off and ICA was closed. The monofilament suture was inserted from ECA and advanced into ICA. 2 hours after surgery, 24 hours reperfusion was performed by removing the monofilament suture. Sham rats were subjected to the same procedure without MCAO/R.

Neurological function test

Neurological function was assessed on modified Neurological Severity Scores (mNSS) [43] and Morris water maze [44] on post-MCAO/R days 1, 3, 7, 14, 21 and 28 days. The mNSS is a composite test of motor, sensory systems, reflexes and balance, the scoring method follows the description [43]. The higher scores indicate more severe neurological impairment of CNS.

To assess the spatial learning and memory of rats, we performed Morris water maze (MWZ) test. The MWZ pool (140 cm diameter, 60 cm height) were split into four quadrants with a platform (9 cm diameter, 23 cm height), which was placed in the third quadrant. MWZ test were divide to three parts, visible platform test from day 1 to day 2, hidden platform test from day 3 to day 7, probe test on day 8. In visible platform test, rats were given 20 seconds to acclimatize on the platform and then placed in each quadrant separately and had 60 seconds to reach the platform. In hidden platform test, the platform was hidden 1 cm under water, and then rats were placed in each quadrant separately and had 60 seconds to reach the platform. In probe test, the platform was removed and the rats were placed to each quadrant for 60 seconds. The time and movement track of rats in MWZ test was recorded by Ethovision XT monitoring system (Noldus, China). The escape latency, swimming path, and target zone frequency were analyzed.

Neurological regeneration test

Neurological regeneration was tested by Nissl staining and Luxol fast blue staining [45] according to the instruction of manufacture. Rats were anesthetized by 3% pentobarbital sodium (42 mg/kg), injection and intracardially perfused with paraformaldehyde. The brain tissues were isolated and fixed in 4% paraformaldehyde for 24 hours, after which the hippocampus and cerebral cortex of the brain tissue were isolated. The brain tissues were dehydrated with 30% sucrose solution at 4°C for 3–5 days. After that, the brain tissues were preserved in optimum cutting temperature compound (OCT) for 6 hours at room temperature. Then the brain tissues were sliced into 8 and 20 μm thick sections at −20°C, respectively. Luxol fast blue (LFB) staining can
demonstrated myelin. Briefly, 8 μm coronal brain slices were placed in LFB solution (Solarbio, China) at 60°C for 4 hours, then transferred to Lithium Carbonate for 30 seconds, obtained the image under bright field microscope (Leica, Germany). Nissl staining can demonstrate Nissl bodies. Briefly, 20 μm coronal brain slices were dehydrated in graded alcohol, staining by 0.1% cresyl violet (Solarbio, China), dehydrated in graded alcohol and xylene again, then transfer to Nissl Differentiation Solution (Solarbio, China) for 30 seconds, obtained the image under bright field microscope (Leica, Germany).

**Enzyme linked immunosorbent assay (ELISA)**

The secretion of pro-inflammatory cytokines IL–1β and IL-18 in cells and brain tissue were detected by ELISA kits (abcam, China) [46], respectively. As manufacturer’s instruction, collected the cells and the cells were centrifuged at 2000 g for 10 minutes, after that, the cell supernatant was collected. Brain tissue lysates were centrifuged at 18,000 g for 20 minutes, after that the supernatants were collected. 50 μL per well samples supernatant and 50 μL per well antibody cocktail were added in to 96 plate wells, and the incubated for 1 hours at room temperature sharked with 400 rpm. After washed with 1X Wash Buffer PT, 100 μL 3,3,5,5-tetramethylbenzidine (TMB) development solutions were added in each well and incubated for 10 minutes in the dark sharked with 400 rpm. 100 μL of stop solution were added in each well, sharked whit 1 min for mixed and then the secretion of IL-1β and IL-18 will determine by spectrometry at 450 nm after incubation.

**Infarct volume assessment**

The infarct volume was analyzed by 2,3,5-triphenyltetrazolium chloride (TTC, Sigma China) staining [47]. After model establishment, rats were euthanized immediately after reperfusion, then brain was harvested and frozen for 5 minutes at −80°C. Brains were cut into 2 mm sections placed in 2% TTC solution for 30 minutes at 37°C in dark. The stained brain slices were arranged in the order, and then labeled, ruled and photographed, and the infarct volume was calculated using Image-Pro Plus image processing software (infarct volume = infarct area of each slice × 2 mm).

**Western blot assay**

Rats were euthanized followed the guideline of Kunming medical university, and the brain tissue was isolated. The total protein of cells or brains tissue were extracted by RIPA buffer (abcam, China) with protease inhibitor cocktail (abcam, China). Pierce BCA protein assay (abcam, China) were performed to determine protein concentration. The protein sample were separated by sodium dodecyl sulfate-12 polyvinylidene gel (SDS-PAGE) electrophoresis and then transferred to polyvinylidene fluoride (PVDF) membranes [48]. The membranes were blocked with 5% nonfat milk in TBST for 1 hour at room temperature. Membranes were incubated with primary antibodies caspase-1 (1:100, ab74297, abcam), IL-1β (1:1000, ab205924, abcam), NLRP3 (1:1000, ab263899, abcam), gasdermin (1:1000, ab219800, abcam), cleaved N-terminal GSDMD (1:1000, ab215203, abcam) overnight at 4°C, respectively. The membranes were washed with TBST, then incubated with secondary antibody for 2 hours at room temperature. Membranes were visualized by enhanced chemoluminescence (ECL) system. GADPH was used as a loading control. The protein bands intensities were imaged using an Odyssey Fc Image System and protein intensities were normalized GADPH.

**Statistics**

All data were performed using GraphPad Prism 9.0 and expressed as mean ± standard deviation (± SD), differences between two groups were analyzed using Student’s t-test, differences between three or more groups were analyzed by one-way ANOVA with a Tukey test. Differences with p < 0.05 were considered a significantly differences [18].
Results

ZEB2 prompted astroglrosis in vitro under OGD/R

In this manuscript, we assumed that ZEB2 promoted astroglrosis and therefore promote neuron regeneration via decreased neuron pyroptosis. Therefore, we first verified the function of ZEB2 in inducing astroglrosis. Astroglrosis is accompanied by increased glial fibrillary acidic protein (GFAP) expression, thus, elevated expression of GFAP is a hallmark signal for activation of astrocyte [28,49].

To evaluate expression of ZEB2 in astrocyte under OGD/R condition, Western blot assay was performed. The results demonstrated that protein expression of ZEB2 was significantly increased in OGD/R group compared with Ctrl group, and the expression of ZEB2 was further increased when transfected with pcDNA3.1 ZEB2, which indicated that the transfection of pcDNA3.1 ZEB2 was successful (Figure 1a, b). To further evaluate the subcellular localization and expression of ZEB2 in astrocytes, immunofluorescence assay was performed. As shown in Figure 1c-E, the astrocyte marker GFAP is mainly expressed in the cytoplasm, while ZEB2 is mainly expressed in the nucleus of astrocytes. In addition, the expression of ZEB2 and GFAP was significantly upregulated in OGD/R group compared to the control group, while a further increase of GFAP expression was observed when pcDNA3.1 ZEB2 was transfected. This result indicated that OGD/R induced astroglrosis, and the upregulation of ZEB2 further increase the activated astrocytes.

ZEB2 protect neuron from pyroptosis via prompting astroglrosis

Prevailing evidence suggested that ZEB2 mainly expressed in astrocytes of brain tissue in rats, and knockdown of ZEB2 in vivo suppressed neural regeneration [28]. To investigate the relationship between ZEB2 expression in astroglrosis and neural regeneration in vitro, we co-cultured astrocytes and neurons. The expression of GFAP and ZEB2 in co-cultured astrocytes were detected by western bolt, the results shown that the expression of ZEB2 and GFAP were increased under OGD/R condition, and the upregulation of ZEB2 further increased the expression of GFAP (Figure 2a-c).

To investigate the role of astrocytes-derived ZEB2 exerts in neuron regeneration, immunofluorescence assay was performed on co-cultured neurons. Brain-derived neurotrophic factor (BDNF) is a member of neurotrophic, it promoted new neurons growth and differentiation [50]. A specific marker of neuron Microtubule-associated proteins 2 (MAP2) was

![Figure 1](image-url)

**Figure 1.** ZEB2 promoted the activation of astrocyte after OGD/R in vitro. A: Western blot assay of ZEB2 expression in astrocyte. B: Quantification of Western blot band intensity. C: Immunofluorescence staining for GFAP (green fluorescence), ZEB2 (red fluorescence), and DAPI (blue) in astrocytes. D: Fluorescence intensity of ZEB2. E: Fluorescence intensity of GFAP. (Error bars represent mean ± SD, Magnification:400×; *p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001).
used to detect the number of neurons. The expression of BDNF and the number of neurons were both decreased under OGD/R condition, and overexpression of ZEB2 in astrocytes attenuated this trend (Figure 3a-c). Then, to detect the proliferation of neurons, the expression of Brdu in neurons was detected. Results demonstrated that the expression of Brdu, was decreased under OGD/R condition. While overexpression of ZEB2 in astrocytes reversed this trend, which indicated the increased proliferation of neurons (Figure 3d-f). In addition, proliferation and apoptosis of neuron was detected by CCK8 and flow cytometry, respectively. Results of CCK8 assay demonstrated that proliferation of neuron was

Figure 2. ZEB2 promoted astrogliosis after OGD/R in vitro. A: Western blot assay of ZEB2 and GFAP expression in astrocyte. B: Quantification of ZEB2 Western blot band intensity. C: Quantification of ZEB2 Western blot band intensity. (Error bars represent mean ± SD; *p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001).

Figure 3. Astroglisis promoted neuron regeneration in vitro. A: Immunofluorescence staining for NeuN+ (green fluorescence), BrdU + (red fluorescence), and DAPI (blue fluorescence) in neurons. B: Immunofluorescence staining for BDNF (green fluorescence), MAP2 + (red fluorescence) and DAPI (blue fluorescence) in neurons. C: Fluorescence intensity of NeuN. D: Fluorescence intensity of BDNF. E: Fluorescence intensity of BrdU. F: Fluorescence intensity of MAP2. G-H: Results of flow cytometry demonstrating astrogliosis alleviated apoptosis of neuron. I: Results of CCK8 demonstrating astrogliosis promoted proliferation of neuron (Error bars represent mean ± SD, Magnification:400×; *p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001).
decreased in OGD/R group compared with Ctrl group, while this trend alleviated by up-regulated expression of ZEB2 in astrocytes (Figure 3i). The apoptosis of neuron shown the opposite result (Figure 3g-h). These results indicated that, astrogliosis and neuron damage were observed under OGD/R condition, while up-regulated ZEB2 expression promoted a higher degree of astrogliosis, which promoted neuron regeneration.

Previous studies reported that inflammasome-mediated pyroptosis play a key role in I/R injury of brain [51]. It is reasonable to assume that higher degree of astrogliosis induced by ZEB2 can protect neural regeneration from inflammation and pyroptosis. The results demonstrated that the expression of pyroptosis markers, i.e., caspase-1, IL-1β, NLRP3, GSDMD and cleaved N-terminal GSDMD was increased under OGD/R condition, and the overexpression of ZEB2 in astrocytes reversed the trend (Figure 4a-f). The results of ELISA shown that the secretion of IL-1β and IL-18 was increased under OGD/R condition, the overexpression of ZEB2 in astrocytes suppressed the secretion of these inflammatory cytokines (Figure 4g-h). Taken together, our results demonstrated that ZEB2 promoted astrogliosis, which protect neuron from pyroptosis under OGD/R condition.

**ZEB2 promoted neural regeneration in vivo**

Then, we extended our work into an in vivo model. AAV-ZEB2 were injected into the brain at 20 days prior to middle cerebral artery occlusion/reperfusion (MCAO/R) model established, the experimental produces were shown in Figure 5. We first detected the expression of ZEB2 in brain tissue of rats in different groups, and found that ZEB2 was aberrantly overexpressed in MACO/R model. The injection of AAV further increased the expression of ZEB2 in brain tissue. Which suggested the successful infection of rAAV. (Figure 6a-b). Morris water maze test was performed to evaluate the spatial learning and memory capacity of rats. The MCAO/R rats needed longer time and more complicated route to find the hidden platform than sham group, and the overexpression of ZEB2 accelerated the time for MCAO/R rats finding the hidden platform, which also significantly simplified the route (Figure 7a-b). Moreover, the probe test shown the time in the target quadrant in MCAO/R rats were significantly decreased compared with Sham rats, and the overexpression of ZEB2 attenuated the trend (Figure 7c). Similarly, the score of the modified Neurological Severity Scores (mNSS) was

![Figure 4](image-url)

**Figure 4.** Astrogliosis promoted neuron regeneration by decreasing pyroptosis in vitro. A: Western blot assay of pyroptosis relative protein expression in neuron. B: Quantification of Caspase-1 Western blot band intensity. C: Quantification of IL-1β Western blot band intensity. D: Quantification of NLRP3 Western blot band intensity. E: Quantification of GSDMD Western blot band intensity. F: Quantification of cleaved N-terminal GSDMD Western blot band intensity. G: ELISA analysis of the secretion of IL-1β in neuron. H: ELISA analysis of the secretion of IL-18 in neuron. (Error bars represent mean ± SD; *p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001).
significantly increased in MCAO/R group compared with sham group (Figure 7d). However, mNSS score was decreased when ZEB2 was overexpressed compared with MCAO/R group. TTC staining was used to detect the infarct area (Figure 7e). It indicated that the injection of AAV-ZEB2 alleviated the tissue loss caused by ischemia. These results suggested that MCAO/R rats had a neurological deficit, and ZEB2 overexpression alleviated this deficit. Nissl body is a large granular body in neuron, which is a kind of rough endoplasmic reticulum with rosettes of free ribosomes, and is the site of protein synthesis [52]. Nissl body decreased in injury neuron [53]. The results of Nissl staining (Figure 7f) indicated that the number of Nissl bodies was decreased in cerebral cortex, hypothalamus and hippocampus in MCAO/R rats compared with sham rats, while ZEB2 reversed the result. The above experiments demonstrated that up-regulated ZEB2 promoted neuron regeneration after MCAO/R injury. To detect the role of ZEB2 on the neuronal damage, Luxol Fast Blue (LFB) staining were performed to observe the myelin and Bielschowsky sliver staining was used to observe axons. The number of myelin and axon was significantly decreased in cerebral cortex, hippocampus and hypothalamus of MCAO/R rats compared with sham rats, and then ZEB2 reversed the result (Figure 7g). Also, demyelination in nuclei observed in MCAO/R rats was also alleviated by ZEB2 overexpression. To further investigated the mechanism of ZEB2 attenuated neuron damage in MCAO/R rats, the level of inflammasome and pyroptosis was detected by ELISA and Western blot. (Figure 7h-o). We found upregulated ZEB2 alleviate the protein expression of pyroptosis markers, i.e., caspase-1, IL-1β, IL-18, NLRP3, GSDMD and cleaved N-terminal GSDMD. Taken together, our results demonstrated that ZEB2 promoted neural regeneration of brain in MCAO/R rats by decreased the inflammation and pyroptosis.

Discussion
Cerebral ischemia causes irreversible cell death or impairment of neuron and astrocytes, which
Figure 7. Overexpression of ZEB2 in brain tissues of MCAO/R rats promoted neuron regeneration by alleviating pyroptosis. A: Representative track images of each group mice. B: Mean escape latency time during the orientation navigation test on 1, 3, 7, 14, 21, 28 days after MCAO/R. C: Time of the rats stay in the target quadrant on 28 days after MCAO/R. D: mNSS score on 1, 3, 7, 14, 21, 28 days after MCAO/R. E: Representative images of brain slices stained by TTC in different groups at 28 days after MCAO/R. F: Staining by Nissl showed the Nissl bodies of hippocampus and cerebral cortex. G: Staining by Luxol fast blue (LFB) showed the myelin of hippocampus and cerebral cortex. H: Western blot analysis of the expression of pyroptosis protein i.e., caspase-1, IL-1β, NLRP3, gasdermin, cleaved N-terminal GSDMD in brain tissue. I: Relative protein expression of caspase-1. J: Relative protein expression of IL-1β. K: Relative protein expression of NLRP3. L: Relative protein expression of gasdermin. M: Relative protein expression of cleaved N-terminal GSDMD. N: ELISA analysis of the expression of IL-1β in brain tissue of rats on 28 days after MCAO/R. O: ELISA analysis of the expression of IL-18 in brain tissue of rats on 28 days after MCAO/R. (Error bars represent mean ± SD, Magnification:400×; *p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001.)
factors

**Conclusion**

Our findings demonstrated that ZEB2 promoted astrogliosis, which therefore enhances neural regeneration by decreasing the inflammation and pyroptosis. It offered a new potential therapeutic method for cerebral I/R injury.

**Disclosure statement**

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Availability of Data and Materials

All data and materials are available from the corresponding author on reasonable request.

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