ORIGINAL RESEARCH Hypoxia Inducible Factor-I α Through ROS/NLRP3 Pathway Regulates the Mechanism of Acute Ischemic Stroke Microglia Scorching Mechanism

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Purpose: In vitro experiments explored how the hypoxia-induced factor-1 α (HIF-1 α) regulates the regulation of pyroptosis in microglial cells (BV 2) in acute ischemic stroke through ROS/NLRP3 pathway.

Methods: The microglia acute phase oxygen-glucose deprivation/reoxygenation (OGD/R) was established, CCK-8 was applied to determine the optimal timing of intervention modeling. HIF-1a was overexpressed with stabilizer GF-4592 and HIF-1a small molecule interfering RNA (HIF-1a-siRNA), which was divided into group A (blank group), group B (OGD/R model group), group C (model +FG-4592 intervention group), group D (model+siRNA negative control group) and group E (model+HIF-1α-siRNA group). Cell proliferation of different groups was measured by CCK-8 assay. Pyroptosis and intracellular ROS levels were measured by flow cell technology. IL-18, IL-1β levels were measured by ELISA. HIF-1α, GSDMD-D, GSDMD-N, clean-Caspase-1 and NLRP3 protein expression levels were measured by Western blot. On the above experiments, ROS and NLRP3 response experiments were performed to explore how HIF-1a regulates pyroptosis through ROS/NLRP3 pathway.

Results: Hypoxia for 6 h then reoxygenation for 12 h was the optimal intervention time. Compared with groups B and D, cell proliferation in group C was significantly enhanced, pyroptosis, intracellular levels of ROS, IL-18, IL-18 and the expression of GSDMD-D, GSDMD-N, clean-Caspase-1, and NLRP3 proteins were significantly decreased in group C (P < 0.05). However, in group E, the performance of these test indicators were exactly the opposite, and the difference was statistically significant (P < 0.05). Through ROS and NLRP3 response experiments, it was found that HIF-1a Inhibition of Pyroptosis by inhibiting ROS/NLRP3 pathway.

Conclusion: Overexpression of HIF-1a factor can inhibit microglia pyroptosis. HIF-1a factor has an inhibitory effect on the ROS/ NLRP 3 pathway, which can inhibit the pyroptotic process in microglia.

Keywords: hypoxia-inducible factor-1a, acute ischemic stroke, microglial pyroptosis, reactive oxygen species, nucleotide-binding oligomerization domain-like receptor protein 3

Introduction

Ischemic stroke is a disease with a high fatality rate. Due to its poor prognosis, it puts great pressure on global health management.¹ Ischemic stroke in the acute phase involves pale tissue with mild enlargement of brain tissue in the hypoxic cell necrotic area between 1 and 24 hours after onset, and significant changes in both microglia and endothelial cells.² The hypoxia-inducible factor-1 α (HIF-1 α) is an important transcription factor regulating the complex cellular stress response of the body.³ The HIF-1 α factor plays an important role in regulating the intracellular environment in the ischemic and hypoxic environment and activating the transcription of various downstream target genes.^{4,5}

Microglia are the earliest and fastest innate immune cells with stress response after brain tissue injury, and researchers pay attention to the inflammatory response of ischemic stroke.^{6–8}

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In the early stages of brain injury, microglia undergo rapid activation, a process that leads to the release of various cytokines, which in turn causes persistent inflammatory damage to the brain tissue.^{9–11} Pyroptosis, a special form of cell death, is a manifestation of cell swelling with condensed chromatin reaches plasma membrane permeability, subsequent lysis of swollen cells releases inflammatory factors involved in the inflammatory response after brain injury.^{12–14} Microglial pyroptosis is closely related to the inflammatory cascade of ischemic stroke. The nucleotide-binding oligomerization domain-like receptor protein 3 (NLRP3) formed by microglial activation participates in the occurrence of pyroptosis and constitutes the classical pyroptosis pathway.^{15,16}

Therefore, this study intends to deeply explore the regulation mechanism of HIF-1 α on microglial pyroptosis in acute cerebral ischemia and hypoxia through in vitro cell modeling, which will help to deepen the clinical understanding of acute ischemic stroke and provide potential targets for clinicians in the precise treatment of acute ischemic stroke.

Materials and Methods

Material

BV2 Mouse Microglia Cell Strain

BV 2 mouse microglia cells were obtained from Wuhan Prosai Biotechnology Co., LTD. (stock number: CM-0493), and the cells were introduced to the library for less than 5 generations.

2. Reagents and consumables

2.1 Reagents and consumables are detailed in Table 1.

2.2 Details of the experimental instruments are shown in Table 2.

Table I Reagent Schedule

Reagent/Consumable	Manufacturer	Art.No
Specific culture medium for the BV 2 cells	Purnosa	CM-0493
DMEM (sugar-free) medium	GIBCO	11966-025
Pancreatin (0.25% Trypsin-EDTA)	GIBCO	25200-056
PBS phosphate buffer powder	Beijing Zhongshan Jinqiao Biology	ZLI-9062
The CCK-8 cell proliferation/toxicity assay kit	All-style gold creatures	FC101-03
DMSO	SIGMA	D2650
Lactate dehydrogenase (LDH) kit	Nanjing completed	A020-2-2
Block-iT Alexa Fluor Red	Thermo Fisher	14750-100
lipofectamin RNAiMAX	Thermo Fisher	13778-150
siRNA Negative control	Jimma gene	
siRNA-HIF-1α -835	Jimma gene	
siRNA-HIF-1α -1235	Jimma gene	
siRNA-HIF-1a -1998	Jimma gene	
Reactive oxygen species (ROS) assay kit	Nanjing completed	E004-1-1
Mouse IL-1B ELISA Kit	Lianke biology	EK201B/3-48
Mouse IL-18 ELISA Kit	Lianke biology	EK218-48
Annexin V PE/7AAD Kit	BD	559763
25 cm ² culture bottle	Corning	430639
96 Hole plate	Corning	3599
6 Hole plate	Corning	3516
TRIzoI™ Reagent	ambion	15596026
5X All-In-One RT MasterMix	abm	G492
EvaGreen Express 2×qPCR MasterMix-Low Rox	abm	MasterMix-EL
Easy II Protein Quantitative Kit (BCA)	All type of gold	DQ111-01
RIPA lysate	Dr De	AR0105
Protease inhibitor	Dr De	AR1178

(Continued)

Reagent/Consumable	Manufacturer	Art.No
Goat Anti-Rabbit IgG H & L (HRP)	Abcam	ab205718
Goat Anti-Mouse IgG H & L (HRP)	Abcam	ab205719
Protein was prestained with Marker	Solaibao	PR1910
PVDF Transfer Membrane (0.45 μm)	Millipore	IPVH00010
PVDF Transfer Membrane (0.22 μm)	Millipore	ISEQ00010
SuperSignal™ West Pico PLUS Chemiluminescent Substrate	Thermo Fisher	34580
Beta-Actin Loading Control antibody Mouse MAb	Yiqiao Shenzhou	100166-MM10
Anti-HIF-1 alpha antibody [EPR16897]	Abcam	ab1 7948 3
GSDMDC1 Antibody (H-11)	SCBT	sc-393581
Anti-NLRP 3 antibody [EPR23094-1]	Abcam	ab263899
Anti-pro Caspase-1 + p10 + p12 antibody [EPR16883]	Abcam	ab179515

Table 2 Experimental Instruments

Instrument	Manufacturer	Model	
CO ₂ Cell incubator	Shanghai Likang Instrument Co., LTD	Smart Cell HF-90	
Biosafety cabinet	Shanghai Likang Instrument Co., LTD	HFI200LC	
Desktop low speed centrifuge	Shanghai Flying Pigeon Instrument Co., LTD	DK-80	
Thermostat water bath	Shanghai Jinghong Instrument Factory	TGL-16GB	
-20°C cryogenic refrigerator	Hefei Meiling Company	BCD-249LCK	
-80°C cryogenic refrigerator	Hefei Meiling Company	DW-HL388	
Manual, single-channel pipette	Eppendorf	Research plus	
Nitrogen canister	Chengdu Jinfeng Instrument Factory	YDS-50-125	
Flow cytometry	BD	LSRFortessa	
ELIASA	The Bio-Rad Corp	xMarkTM	
The BG-Power600i conventional electrophoresis	Beijing Baijing Biotechnology Co., LTD	BG-Power600i	
apparatus power supply			
Electrophoresis bath	Beijing Liuyi Instrument Factory	DYCZ-21	
Gel imaging system	Shanghai day can	2500	
PCR appearance	Bio-Rad	MyCycler Thermal Cycler	
Real Time PCR instrument	ABI	7500 Fast	
Chemiluminescence imager system	Shanghai Qinxiang Scientific Instrument Co., LTD	Chemiscope 3000	
Protein transmembrane instrument	Bio-Rad	Mini-PROTEAN Tetra system	
Electrophoresis apparatus trophoresis	Beijing Liuyi	DYCZ-24DN	

Method

1. Cell line cultivation Cell culture conditions were MEM medium + 10% FBS + 1%PS, 37° C, 5% CO₂ with saturated humidity. The cells were grown continuously for 8 days and detected the OD values by a microplate reader every 24 hours.

2. In vitro experiments

2.1 To model oxygen-glucose deprivation reoxygenation in the acute phase (OGD/R)

After 24 h incubation, BV 2 cells were replaced. Cells were placed in a 37°C saturated gas incubator with 95% nitrogen plus 5% oxygen for 3 h and 6 h of hypoxia culture and replaced with fresh MEM complete medium, and cells were replaced into a CO_2 incubator for normal incubation for 12 hours. The control group was always cultured in the CO_2 incubator for the same time.

2.2 CCK-8 detects cell viability to determine the optimal timing of intervention

3. Investigate the effect of HIF-1a on cell pyroptosis in the OGD/R model

3.1 Experimental grouping

(1) Blank group: The BV 2 cells were routinely cultured.

(2) OGD/R model group: BV2 cells were subjected to hypoxia for 6 hours in sugar-free basal medium, followed by reoxygenation for 12 hours in normal medium.

(3) OGD/R model + FG-4592 intervention group: After BV2 cells were cultured for a certain period of time, FG-4592 (50 μ M) was added intervention for 1 hour, hypoxia for 6 hours in a sugar-free basal medium, followed by reoxygenation in a normal medium for 12 hours.

(4) Model + siRNA negative control group: After transfection of siRNA NC into BV2 cells for 20 hours, hypoxia was observed in sugar-free basal medium for 6 hours, followed by reoxygenation in normal medium for 12 hours.

(5) Model + HIF-1 α -siRNA group: Transfection of siRNA-HIF-1 α -835 into BV2 cells for 20 hours, hypoxia was observed in sugar-free basal medium for 6 hours, followed by reoxygenation in normal medium for 12 hours.

3.2 Cell proliferation and pyroptosis were detected in the different groups

Different grouped cell proliferation was assessed by CCK-8. Cell pyroptosis was determined in different groupings by flow cytometry.

3.3 Detection of intracellular reactive oxygen species (ROS) levels by flow cytometry

3.4 The level of IL-1 β , IL-18 concentration in the cell supernatants was determined by ELISA

3.5 Western Blot detected the expression of HIF-1α, GSDMD-D, GSDMD-N, cle-Caspase-1, and NLRP 3 proteins in different groups

3.6 On the above experimental basis further explored the role of ROS in HIF-1 α regulation of microglia pyroptosis

Using ROS inhibitor N-acetylcysteine (NAC) to further explore the role of ROS in HIF-1 α Regulation of cell apoptosis. The ROS inhibitor NAC combined with HIF-1 α intervention could be divided into blank group, blank + ROS + NAC group, OGD/R model + FG-4592 intervention group, OGD/R model + FG-4592 intervention+ROS+NAC group, model + HIF-1 α -siRNA group, and model+HIF-1 α -siRNA+ROS+NAC group. Intracellular ROS activity, NLRP3 protein expression, LDH activity, IL-1 β , and IL-18 were examined again.

3.7 On the above experimental basis, we further explored the role of NLRP3 protein in the regulation of HIF-1 α in microglia pyroptosis

In this study, the NLRP3 inhibitor MCC950 was used to further validate the role of NLRP3 in regulating cell pyroptosis by HIF-1 α . The NLRP3 inhibitor MCC950 combined with HIF-1 α intervention could be divided into blank, blank +NLRP3+MCC950, OGD/R+FG-4592 intervention group, OGD/R + FG-4592 intervention + NLRP3 + MCC950, model + HIF-1 α -siRNA, and model+HIF-1 α -siRNA+NLRP3+MCC950 group. Intracellular ROS activity, NLRP3 protein expression, LDH activity, IL-1 β , and IL-18 were then again examined.

Statistical Analysis

All data were represented by mean \pm standard deviation ($\bar{x} \pm S$). Spss19.0 software was used to analyze the data of each group. One-way ANOVA was used to analyze the data between groups, *P*<0.05 showed significant differences.

Results

The BV 2 Cell Growth Curve

Cultivate BV2 cells in vitro, detect OD values at the wavelength of 450 nm using an enzyme-linked immunosorbent assay every 24 hours, and plot growth curves. From the growth curve, it can be seen that cells are in a slow growth latent period from 1 to 3 days, and the slope of the growth curve is relatively large from 3 to 5 days. The curve of slow growth from 5 to 7 days presents a plateau like flat top period (Figure 1).



Figure I BV 2 cell growth curves. (A) Growth plot of BV 2 cells. (BI-B7) Plot of BV 2 cell proliferation at days I-7.

Determine the Optimal Intervention Time as 12 Hours of Reoxygenation After 6 Hours of Hypoxia

Compared with the blank group of the BV 2 cell lines, the BV2 cell line showed a decrease in cell viability after 3 hours of hypoxia then 12 hours of reoxygenation, but the difference was not statistically significant (P>0.05). The cell viability of the model significantly decreased after 6 hours of hypoxia then 12 hours of reoxygenation, the difference was statistically significant (P<0.05) (Table 3). The optimal intervention timing was ultimately determined to be 6 hours of hypoxia and 12 hours of reoxygenation.

Cells Proliferation Was Significantly Higher in OGD/R Model + FG-4592 Intervention Group but Significantly Decreased in Model + HIF-1 α -siRNA Group Cells

The analysis of variance showed that compared with the blank group, the cell proliferation in the other four groups was significantly decreased (P<0.05). Compared with the OGD/R model group, cell proliferation was significantly higher in the OGD/R model + FG-4592 intervention group but significantly decreased in the model + HIF-1 α -siRNA group (P<0.05). Compared with the OGD/R model + FG-4592 intervention group, cell proliferation was significantly reduced in the model + siRNA negative control group and model + HIF-1 α -siRNA group (P<0.05). Compared with the model + siRNA negative control group, cell proliferation was significantly reduced in the model + siRNA negative control group, cell proliferation was significantly reduced in the model + HIF-1 α -siRNA group (P<0.05). (Table 4).

The OGD/R model+FG-4592 Intervention Group Significantly Reduced the Cell Apoptosis Rate, While the Cell Death Rate Significantly Increased in the model+HIF-I α -siRNA Group

Analysis of variance was used. Compared with the OGD/R model group, the OGD/R model+FG-4592 intervention group significantly reduced the cell apoptosis rate, while the cell death rate significantly increased in the model+HIF-1 α -siRNA

6 Hours ($x \pm 5$, n=5)						
Experimental Grouping	OD Value					
	Hypoxic for 3 h by Reoxygenation for 12 h	Hypoxic for 6 h by Reoxygenation for 12 h				
Blank group	0.346±0.038	0.373±0.030				
The OGD/R model group	0.296±0.038	0.197±0.043				
t	2.106	5.307				

0.05

Table 3 Results of BV 2 Cell Proliferation Detection in Two Groups Under Hypoxia for 3 Hours and 6 Hours ($\bar{x} \pm S$, n=5)

Notes: Compared with the blank group, *P <0.05, was statistically significant.

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0.001*

Table	4	Results	of	ΒV	2	Cell	Proliferation	Detect	ion	in	5	Different
Experi	me	ntal Gro	ups	; (x :	± 2	S, n=!	5)					

Experimental Grouping	OD value
Blank group	0.702±0.045
The OGD/R model group	0.402±0.039 [∆]
The OGD/R model + FG-4592 intervention group	0.511±0.058 [∆] ▲
Model + siRNA negative control group	0.380±0.054 ^{∆▽}
Model + HIF-1 α -siRNA group	0.291±0.030 ^{△▲▽▼}

Notes: P <0.05 for $^{\triangle}$ and blank group; P <0.05 for $^{\clubsuit}$ and OGD/R model group; P <0.05 for $^{\bigtriangledown}$ and OGD/R model + FG-4592; P <0.05 for $^{\checkmark}$ and model + siRNA negative control group.

group (P < 0.05). Compared with the OGD/R model + FG-4592 intervention group, the cell pyroptosis rate was significantly increased in the model + siRNA negative control group and model + HIF-1 α -siRNA group (P < 0.05). Compared with the model + siRNA negative control group, the cell pyroptosis rate was significantly increased in the model + HIF-1 α -siRNA group (P < 0.05) (Figure 2).

The ROS Levels in OGD/R model+FG-4592 Intervention Group Was Significantly Decreased While the model+HIF-Iα-siRNA Group Was Significantly Elevated

Analysis of variance: compared with the blank group, the ROS levels of the other four groups were significantly increased (P<0.05). Compared with the OGD/R model group, the ROS levels in OGD/R model+FG-4592 intervention group were significantly decreased, while the model+HIF-1 α -siRNA group was significantly elevated (P<0.05). Compared with the OGD/R model + FG-4592 intervention group, the ROS levels were significantly increased in the model + siRNA negative control group and the model + HIF-1 α -siRNA group (P<0.05). Compared with the model + siRNA negative control group, the ROS levels were significantly increased in the model + siRNA negative control group, the ROS levels were significantly increased in the model + HIF-1 α -siRNA group (P<0.05). Compared with the model + siRNA negative control group, the ROS levels were significantly increased in the model + HIF-1 α -siRNA group (P<0.05). (Figure 3).



Figure 2 Pyroptosis was determined by flow cytometry. (A) Comparison of pyroptosis analysis, (B–F): flow cytometry results in different groups. Note: ***P < 0.05 was statistically significant.



Figure 3 Results of ROS levels in BV 2 cells. (A) Different groups' cell ROS bar graph, (B1–B5) ROS results by flow cytometry in different groups. Note: **P < 0.05 was statistically significant.

The Contents of IL-1 β and IL-18 in the OGD/R Model + FG-4592 Intervention Group Was Significantly Reduced While the Model + HIF-1 α -siRNA Group Was Significantly Increased

Compared with the blank group, IL-1 β and IL-18 were significantly increased in the other four groups (*P*<0.05). Compared with the OGD/R model group, the contents of IL-1 β and IL-18 in the OGD/R model + FG-4592 intervention group were significantly reduced while model + HIF-1 α -siRNA group's significantly increased (*P*<0.05). Compared with the OGD/R model + FG-4592 intervention group, IL-1 β and IL-18 were significantly increased in the model + siRNA negative control and model + HIF-1 α -siRNA group (*P*<0.05). Compared with the model + siRNA negative control group, IL-1 β and IL-18 content increased significantly in the model + HIF-1 α -siRNA group (*P*<0.05) (Table 5).

The Expression of the Target Proteins Were Significantly Decreased in the OGD/R Model + FG-4592 Intervention Group While Significantly Increased in He Model + HIF-1 α -siRNA Group

Comparison of expression levels of different grouped proteins after intervention HIF-1 α factor, Compared with the blank group, the expression of the target proteins GSDMD-D, GSDMD-N, cle-Caspase-1, and NLRP3 was significantly increased in the other 4 groups (*P*<0.05). Compared with the OGD/R model group, the expressions of GSDMD-D, GSDMD-N, cle-Caspase-1, and NLRP3 proteins were significantly lower in the OGD/R model + FG-4592 intervention

Experimental Grouping	IL-Iβ (pg/mL)	IL-18 (pg/mL)
Blank group	19.422±4.039	23.055±6.567
The OGD/R model group	36.339±2.886 [∆]	64.060±6.247 [∆]
The OGD/R model + FG-4592 intervention group	23.619±3.653▲	31.335±2.598▲
Model + siRNA negative control group	31.819±0.897 ^{∆▽}	61.533±9.577 ^{∆▽}
Model + HIF-I α -si RNA group	53.501±0.538 ^{△▲▽▼}	95.033±8.858 ^{△▲▽▼}

Table 5 IL-1 β and IL-18 Content in the Supernatant of Cells in Each Group ($\bar{x} \pm S$, n=3)

Notes: Compared with the blank group, $^{\Delta}P$ <0.05; compared with the OGD/R model group, $^{\Delta}P$ <0.05; compared with the OGD/R model + FG-4592 intervention group, $^{\nabla}P$ <0.05; compared with the model + siRNA negative control group, $^{\nabla}P$ <0.05.

group, while significantly higher in the model + HIF-1 α -siRNA group (*P*<0.05). Compared with the OGD/R model + FG-4592 intervention group, the four proteins were significantly increased in the model + siRNA negative control group and the model + HIF-1 α -siRNA group (*P*<0.05). Compared with the model + siRNA negative control group, the four proteins were significantly higher in the model + HIF-1 α -siRNA group (*P*<0.05) (Figure 4).

Inhibition of ROS Would Benefit the Regulation of Cell Pyroptosis by HIF-1 α

Compared with the blank group, the blank + ROS + NAC group significantly decreased ROS activity, NLRP3 protein expression, LDH activity, IL-1 β and IL-18 (*P*<0.05). Compared with the OGD/R model + FG-4592 intervention group, the ROS activity, NLRP3 protein expression, LDH activity, IL-1 β , IL-18 content were significantly reduced in the OGD/R model + FG-4592 intervention + ROS + NAC group (*P*<0.05). Compared with the model + HIF-1 α -siRNA group, the ROS activity, NLRP3 protein expression, LDH activity, IL-1 β and IL-18 content in the model + HIF-1 α -siRNA + ROS + NAC group were significantly reduced (*P*<0.05) (Figure 5).

Inhibition of NLRP3 Would Benefit the Regulation of Cell Pyroptosis by HIF-1 α

Compared with the blank group, the ROS activity, NLRP3 protein expression, LDH activity, IL-1 β and IL-18 were significantly reduced in the blank + NLRP3 + MCC950 group (*P*<0.05). Compared with the OGD/R model + FG-4592 intervention group, the ROS activity, NLRP3 protein expression, LDH activity, IL-1 β and IL-18 content were significantly reduced in the OGD/R model + FG-4592 intervention + NLRP3 + MCC950 group (*P*<0.05). Compared with the model + HIF-1 α -siRNA group, the ROS activity, NLRP3 protein expression, LDH activity, IL-1 β and IL-18 content were significantly decreased in the model + HIF-1 α -siRNA + NLRP3 + MCC950 group (*P*<0.05) (Figure 6).



Figure 4 Protein expression of HIF-1*a*, GSDMD-D, GSDMD-N, cle-Caspase-1, and NLRP3 in the BV 2 cells of each group. (A) Comparison of expression levels of different grouped proteins, (B) Plot of different proteins.

Notes: ${}^{a}P < 0.05$, compared to the blank; ${}^{b}P < 0.05$, compared to the OGD/R model; ${}^{c}P < 0.05$, compared to the OGD/R model + FG-4592; ${}^{d}P < 0.05$, compared to the model + siRNA negative control group.



Figure 5 Content of NLRP3, ROS, LDH, IL-1 β , and IL-18 in different groupings after ROS inhibition. (A) NLRP3 Column analysis of expression in different groups, (B) ROS Column analysis of expression in different groups, (C) LDH Column analysis of expression in different groups, (D) IL-1 β Column analysis of expression in different groups, (E) IL-18 Column analysis of expression in different groups. (E) IL-18 Column analysis of expression in different groups. Note: **P < 0.05 was statistically significant.



Figure 6 Content of NLRP3, ROS, LDH, IL-1 β , and IL-18 in different groupings after NLRP3 inhibition. (**A**) NLRP3 Column analysis of expression in different groups, (**B**) ROS Column analysis of expression in different groups, (**C**) LDH Column analysis of expression in different groups, (**D**) IL-1 β Column analysis of expression in different groups, (**E**) LDH Column analysis of expression in different groups, (**B**) ROS Column analysis of expression in different groups, (**C**) LDH Column analysis of expression in different groups, (**D**) IL-1 β Column analysis of expression in different groups. (**D**) IL-1 β Column analysis of expression in different groups. (**D**) IL-1 β Column analysis of expression in different groups. (**D**) IL-1 β Column analysis of expression in different groups. (**D**) IL-1 β Column analysis of expression in different groups. (**D**) IL-1 β Column analysis of expression in different groups.

Discussion

Acute ischemic stroke (AIS) is a central nervous system cerebral infarction caused by cerebral ischemia and hypoxia caused by cerebral vascular occlusion. It is a kind of central nervous system disease with the highest clinical mortality and disability rate at present.^{17,18} Hypoxia inducible factor-1 α (HIF-1 α) is an important nuclear transcription factor to maintain the survival of tissues and cells and cell homeostasis after the stimulation of hypoxic environment. It is closely related to poor prognosis such as impaired recovery of nerve cells and severe progression of AIS patients.^{19,20}

HIF-1 is made by α , β Heterodimer of two subunits, HIF-1 α can regulate the expression of downstream proteins and regulate cell proliferation, apoptosis, tumor glycolysis and tumor angiogenesis under the stimulation of hypoxic-ischemic environment.^{21–23} HIF-1 α is an important subunit that determines the biological activity of HIF-1, which can induce HIF-1 α in the environment of ischemia and hypoxia then start downstream gene expression.²⁴ Under normal oxygen condition, the expression level is very low and mainly exists in the cytoplasm, but in hypoxia, a large aggregation of HIF-1 α can transfer from the cytoplasm to the nucleus, and the expression level of HIF-1 α increases, with the extension of cell ischemia and hypoxia.^{25,26} HIF-1 α in patients with cerebral hemorrhage, the significantly up-regulated expression has high predictive value for the poor prognosis of patients.²⁷

Abnormal activation of microglia will aggravate the inflammatory response in AIS patients. Microglia are the earliest and fastest innate immune cells after brain injury, and they are also important initiators and participants of other types of glial cells in brain tissue, which lead to persistent inflammatory brain injury.²⁸ Some studies have found that microglia have two-way effects of anti-inflammatory and pro-inflammatory in the process of ischemic stroke.²⁹ When ischemic stroke occurs, microglia are immediately activated and rapidly migrate to the lesion site, while secreting inflammatory cytokines and cytotoxic substances to aggravate brain tissue damage.³⁰ Some small molecule substances produced by activated microglia can protect neurons. Mastering the interaction between neurons and microglia is conducive to clinicians' understanding of the pathological mechanism of nervous system diseases, thus providing a new direction for patients' precise treatment and improving prognosis.^{31–33} Pyroptosis, a newly identified pro-inflammatory disease with a biological characteristic cell death mode unique to cell swelling, plasma swelling cell membrane, inflammatory cytokine secretion, and apoptosis.³⁴ When cell pyroptosis occurs, NLRP3 expression increases sharply along with its activation and converting inactive caspase-1 into catalytically active caspase-1, activated caspase-1 will further cleave and activate Gasdermin domain-containing protein (GSDMD), while activating the expression of inflammatory cytokines IL-18 and IL-18.^{35,36}

This study found that overexpression of HIF-1 α can inhibit the production of reactive oxygen species (ROS) in microglia. A large amount of ROS accumulation can activate the NLRP3 inflammasome and promote the development of the associated inflammatory cascade.³⁷ When the formation of excess NLRP3 inflammasome activates caspase-inflammasome, the initiation of caspase-1 will not only lead to the production of IL-18 and IL-18 inflammatory factors, but also produce pro-inflammatory intracellular substances that promote the breakage and release of the cytoplasmic membrane.³⁸ Our cell in vitro experiments show that overexpression of HIF-1 α in the OGD/R model would enhance microglial proliferation, and Inhibition of HIF-1 α expression would inhibit cell proliferation. Flow cytometry detection of intracellular ROS display, after overexpression of HIF-1 α factor, it was verified that HIF-1 α could inhibit the production of excessive reactive oxygen species. HIF-1 α suppresses mitochondrial oxidation respiration and the generation of reactive oxygen species, thus effectively reducing the ROS content in vivo. The analysis by Western blot experiments showed that the protein content of GSDMD-D, GSDMD-N, cle-Caspase-1 and NLRP3 increased in the OGD/R model. This study showed that the expression of IL-1 β , IL-18, NLRP3, GSDMD, and cle-Caspase-1 were all increased in microglial pyroptosis in the OGD/R model, and this conclusion is consistent with the conclusion of Qianjiang.³⁹ Our experimental results suggest that ROS and NLRP3 either rise or fall synchronously in the presence of HIF-1 α . Overexpression of HIF-1 α will inhibit the increase of ROS in microglia and then inhibit NLRP3 cell pyroptosis. This study found that overexpression of HIF-1 α factor would reduce the production of IL-1 β and IL-18 inflammatory factors in the cells released by microglial rupture.

Based on the above experimental results, we detected ROS activity, NLRP3 protein expression, LDH activity, IL-1 β , and IL-18 concentration again, after ROS was inhibited by single factor under the conditions of HIF-1 α were unchanged. Statistical analysis found that after the condition of HIF-1 α was unchanged and then the inhibition of ROS, the ROS activity, NLRP3 protein expression, LDH activity, IL-1 β and IL-18 concentration in cells were significantly decreased to achieve the effect of reducing pyroptosis of microglia. After this experiment, we only inhibited NLRP3 protein under other conditions

remained unchanged, and found that the ROS activity was reduced but the difference was not statistically significant, while the NLRP3 protein expression, LDH activity, IL-1 β and IL-18 concentrations were reduced and the difference was statistically significant. This response experiment revealed that HIF-1 α regulates microglial pyroptosis through inhibition of ROS and NLRP3 protein expression. Reducing the inflammatory response in stroke by inhibiting NLRP3 inflammatory-mediated microglial pyroptosis may be a new direction for the treatment of ischemic stroke.⁴⁰

In conclusion, our experimental results confirmed that the overexpression of HIF-1 α inhibited the increase of ROS in microglia and thus inhibited the inflammasome NLRP3, and ultimately inhibited microglia pyroptosis through the ROS/ NLRP3 pathway, thus facilitating the treatment of stroke diseases. The conclusion of this study may provide potential targets for the clinical treatment of ischemic stroke in the acute phase.

There are still shortcomings in this study. First, the recovery test was conducted to detect the Pyroptosis index, which did not involve the detection of related proteins. Second, the related in vivo animal experiments should be further carried out to explore and verify.

Conclusion

In conclusion, our study confirmed that overexpression of HIF-1 α factor can effectively inhibit microglial pyroptosis. Secondly, we found that HIF-1 α factor inhibits mitochondrial oxidative respiration and reactive oxygen species generation to inhibit the elevation of ROS in microglia and suppress inflammasome NLRP3, thus indicating that HIF-1 α factor ultimately inhibits microglia pyroptosis through the ROS/NLRP3 pathway. The conclusion of this study may provide potential targets for the clinical treatment of ischemic stroke in the acute phase.

Data Sharing Statement

The study was approved by the Ethics Committee of Urumqi Friendship Hospital. This study is an in vitro cell experiment with no research risk or adverse effects on patients, and does not involve patient information. The data were anonymized or maintained with confidentiality. The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. The study was conducted in accordance with the Declaration of Helsinki.

Author Contributions

All authors made a significant contribution to the work reported, whether that is in the conception, study design, execution, acquisition of data, analysis and interpretation, or in all these areas; took part in drafting, revising or critically reviewing the article; gave final approval of the version to be published; have agreed on the journal to which the article has been submitted; and agree to be accountable for all aspects of the work. All authors contributed equally to this manuscript.

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Disclosure

The authors do not have any commercial or associative interest that represents a conflict of interest in connection with the work submitted.

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