


RESEARCH ARTICLE

Open Access



Regulatory effects of *Trichinella spiralis* serpin-type serine protease inhibitor on endoplasmic reticulum stress and oxidative stress in host intestinal epithelial cells

Jingbo Zhen^{1†}, Lihao Lin^{1†}, Zhixin Li¹, Feng Sun¹, Yang Han¹, Qiankun Li¹, Yuqi Yang¹, Xueting Liu¹, Junchen Yu¹, Qi Zhang¹, Yixin Lu^{1*}  and Caixia Han^{1*}

Abstract

Endoplasmic reticulum stress (ERS) and oxidative stress (OS) are adaptive responses of the body to stressor stimulation. Although it has been verified that *Trichinella spiralis* (*T. spiralis*) can induce ERS and OS in the host, their association is still unclear. Therefore, this study explored whether *T. spiralis*-secreted serpin-type serine protease inhibitor (TsAdSPI) is involved in regulating the relationship between ERS and OS in the host intestine. In this study, mice jejunum and porcine small intestinal epithelial cells (IECs) were detected using qPCR, western blotting, immunohistochemistry (IHC), immunofluorescence (IF), and detection kits. The results showed that ERS- and OS-related indexes changed significantly after TsAdSPI stimulation, and Bip was located in IECs, indicating that TsAdSPI could induce ERS and OS in IECs. After the use of an ERS inhibitor, OS-related indexes were inhibited, suggesting that TsAdSPI-induced OS depends on ERS. When the three ERS signalling pathways, ATF6, IRE1, and PERK, were sequentially suppressed, OS was only regulated by the PERK pathway, and the PERK-eif2 α -CHOP-ERO1 α axis played a key role. Similarly, the expression of ERS-related indexes and the level of intracellular Ca²⁺ were inhibited after adding the OS inhibitor, and the expression of ERS-related indexes decreased significantly after inhibiting calcium transfer. This finding indicated that TsAdSPI-induced OS could affect ERS by promoting Ca²⁺ efflux from the endoplasmic reticulum. The detection of the ERS and OS sequences revealed that OS occurred before ERS. Finally, changes in apoptosis-related indexes were detected, and the results indicated that TsAdSPI-induced ERS and OS could regulate IEC apoptosis. In conclusion, TsAdSPI induced OS after entering IECs, OS promoted ERS by enhancing Ca²⁺ efflux, and ERS subsequently strengthened OS by activating the PERK-eif2 α -CHOP-ERO1 α axis. ERS and OS induced by TsAdSPI synergistically promoted IEC apoptosis. This study provides a foundation for exploring the invasion mechanism of *T. spiralis* and the pathogenesis of host intestinal dysfunction after invasion.

Keywords *Trichinella spiralis*, serine protease inhibitor, endoplasmic reticulum stress, oxidative stress

Handling editor: Vincent Béringue.

[†]Jingbo Zhen and Lihao Lin contributed equally to this work.

*Correspondence:

Yixin Lu

luyixin@neau.edu.cn

Caixia Han

hancaixia@neau.edu.cn

Full list of author information is available at the end of the article



© The Author(s) 2024. **Open Access** This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit <http://creativecommons.org/licenses/by/4.0/>. The Creative Commons Public Domain Dedication waiver (<http://creativecommons.org/publicdomain/zero/1.0/>) applies to the data made available in this article, unless otherwise stated in a credit line to the data.

Introduction

Trichinellosis is a global food-borne parasitic disease caused by the consumption of raw meat containing infectious larvae of *Trichinella spiralis* (*T. spiralis*), seriously endangering human health and leading to tremendous economic losses in animal husbandry [1]. When people or animals are infected with *T. spiralis*, muscle larvae are released into the stomach, migrate into the duodenum, burrow into the intestinal mucosa and develop into adult worms through four moults [2]. Then, female worms and male worms mate in the small intestine and give birth to newborn larvae. This indicates that intestinal invasion is an important stage of *T. spiralis* infection and determines the occurrence and development of trichinellosis [3]. Therefore, it is exceedingly important to study the physiological or pathological changes in the host intestine after *T. spiralis* invasion.

After invasion by many parasites, including *Schistosoma japonicum*, *Clonorchis sinensis*, *Ascaris*, *T. spiralis*, *Haemonchus contortus*, and *Brugia malayi*, serine protease inhibitors (SPIs) affect the physiological processes of the host, such as inflammation, coagulation, complement activation, fibrinolysis, cell development, and programmed cell death of host [4]. Our previous studies confirmed that *T. spiralis* serpin-type SPI (TsAdSPI) is an important SPI derived from muscle larvae and adult worms that can cause autophagy in small intestinal cells, disrupt the intestinal mucosal barrier, and regulate the host immune response [1, 5]. However, whether it is involved in other physiological processes of the intestine needs further research.

Endoplasmic reticulum stress (ERS) and oxidative stress (OS) are adaptive responses of hosts after being stimulated by stressors such as parasites, bacteria, and viruses. Some studies have shown that viruses can promote the host to form a favourable living environment

for self-replication and pathogenicity by regulating the relationship between ERS and OS after infection [6, 7]. Alfredo et al. [8] showed that the host alleviated *hepatitis C virus*-induced damage through the unfolded protein response (UPR) and antioxidant stress, and initiated an immune response to restore cell homeostasis. However, there are no relevant studies on how parasites regulate the association between ERS and OS, although it has been verified that many parasites, including *T. spiralis*, can cause ERS and OS in the host [9–13]. Thus, on the basis of these studies, this study further explored whether TsAdSPI is involved in the regulatory effect of *T. spiralis* on the relationship between host ERS and OS. Our results will provide a reference for the invasion mechanism of *T. spiralis* and the pathogenesis of trichinellosis.

Materials and methods

Animals and cells

BALB/c mice aged 6–8 weeks were purchased from Harbin Medical University. The porcine small intestinal epithelial cell line J2 (IPEC) was generated by Liu Renqiang from Harbin Veterinary Research Institute.

Preparation of protein, agonists, and inhibitors

The pET-30a-TsAdSPI (accession number: EU263307.1) positive expression bacterial solution was purified on a nickel column (Solarbio, China) and renatured. The endotoxin of the recombinant protein was removed using an endotoxin removal kit (Huzhenbio, China). For the dosage and action time of agonists and inhibitors (MedChemExpress, USA), the reagent specifications and related literature were referenced, as shown in Table 1. Due to the absence of a dosage reference for H₂O₂ and EN460, H₂O₂ at dosages of 0, 50, 100, 200, 500, and 1000 μM and EN460 at dosages of 0, 4, 8, 12, 16, and 20 mM were cocultured with IPEC for 1 h, respectively. Then, a

Table 1 Treatment dosage and time of agonists and inhibitors

Reagent	Function	In vivo experiment		In vitro experiment	
		Dosage	Time	Dosage	Time
4-Phenylbutyric acid (4-PBA)	ERS inhibitor	150 mg/kg	Once a day for 7 d	50 μM	1 h
Tunicamycin (Tm)	ERS agonist	1 mg/kg	24 h	1 μg/mL	3 h
<i>N</i> -acetyl-L-cysteine (NAC)	OS inhibitor	150 mg/kg	24 h	1 mM	1 h
Hydrogen peroxide (H ₂ O ₂)	OS agonist	0.74 mM/kg	24 h		1 h
4-(2-Aminoethyl)-benzenesulfonyl fluoride hydrochloride (AEBSF)	ATF6 inhibitor			200 μM	1 h
MKC3946	IRE1α inhibitor			10 μM	1 h
GSK2606414	PERK inhibitor			5 μM	1 h
EN460	ERO1α inhibitor				1 h
2-Aminoethoxydiphenyl borate (2-APB)	Ca ²⁺ release antagonist			1 μM	1 h

CCK-8 cell activity detection kit (Meilunbio, China) was used to determine the optimal interaction concentration.

Preparation of the detection kit and fluorescent probe

Malondialdehyde (MDA) (Nanjing Jiancheng Biology, China), reactive oxygen species (ROS) (Meilunbio, China), and superoxide dismutase (SOD) (Nanjing Jiancheng Biology, China) detection kits were used to measure the MDA and ROS content, and the SOD activity. The Fluo-4 AM fluorescent probe (Meilunbio, China) was used to measure the intracellular Ca^{2+} level. An enzyme-labelled instrument (BioTek, USA) was used to quantify the optical density of MDA and SOD, as well as the fluorescence intensity of ROS and intracellular Ca^{2+} . Cell apoptosis was detected by an Annexin V-FITC apoptosis detection kit (Meilunbio, China) and flow cytometry (FCM) (Sagecreation, China). The mitochondrial membrane potential was detected by a mitochondrial membrane potential detection kit (Solarbio, China), and the fluorescence intensity was observed using a fluorescence microscope (BioTek, USA).

Animal experiments

Sixty-nine BALB/c mice were randomly divided into 23 groups, with 3 mice in each group. Fifteen mice were randomly divided into 5 groups, including the control group, 4-PBA group, Tm group, TsAdSPI group, and TsAdSPI+4-PBA group, to explore the effects of TsAdSPI on ERS and apoptosis in the host intestine by quantitative real-time polymerase chain reaction (qPCR) and western blotting. Moreover, 15 mice were subjected to immunohistochemistry (IHC) experiments. Then, to study the effects of TsAdSPI on OS and apoptosis, 15 mice were randomly divided into the control group, NAC group, H_2O_2 group, TsAdSPI group, and TsAdSPI+NAC group. Finally, 24 mice were randomly divided into 8 groups, namely, the control group, 4-PBA group, NAC group, TsAdSPI group, TsAdSPI+4-PBA group, and TsAdSPI+NAC group, to explore the effects of TsAdSPI-induced ERS on OS and TsAdSPI-induced OS on ERS. The TsAdSPI, agonists, and inhibitors were injected intraperitoneally into the mice. In the TsAdSPI+inhibitor group, TsAdSPI preceded TsAdSPI, and the dose and stimulation time of TsAdSPI were 50 μ g and 24 h, respectively. According to the studies of Xu et al. [3] and Zhen et al. [1], jejunal tissues were selected for subsequent experiments.

Cell experiments

To probe the effects of TsAdSPI on ERS, OS, and apoptosis in IPEC cells, the cells were divided into the PBS group, 4-PBA group, Tm group, NAC group, H_2O_2 group, TsAdSPI group, TsAdSPI+4-PBA group, and

TsAdSPI+NAC group. Then, to explore the interaction between ERS and OS induced by TsAdSPI, the cells were divided into the PBS group, 4-PBA group, NAC group, TsAdSPI group, TsAdSPI+4-PBA group, and TsAdSPI+NAC group. Next, to study the interactions between ERS and OS, the cells were divided into the PBS group, inhibitor group (including AEBSE, MKC3946, GSK2606414, EN460, and 2-APB), TsAdSPI group, and TsAdSPI+inhibitor group. Finally, to study the activation sequence of ERS and OS, the cells were divided into 0, 1, 2, 3, 6, and 9 h groups. In the TsAdSPI+inhibitor group, the IPEC were treated with inhibitor and then stimulated with 10 μ g/mL TsAdSPI for 3 h. The stimulation dosage and time of TsAdSPI were described in a previous study [1].

qPCR

Total RNA was extracted from tissues or cells with a M5 Universal RNA Mini Kit (Mei5bio, China) and reverse transcribed into cDNA using an All-in-One First Strand cDNA Synthesis Kit III (with dsDNase) for qPCR (Seven Biotech, China). qPCR was performed via a Roche LightCycler 480 system (Roche, Switzerland), and the results were calculated using the $2^{-\Delta\Delta Ct}$ method. The primer sequences are shown in Table 2.

Table 2 Primers of the detected genes

Gene	Sequence (5'-3')	Access number	Origin
GAPDH	F: GATTCCACCCACGGCAAGTTCC R: AGCACCAGCATCACCCCATTTG	NM_001206359.1	Swine
GAPDH	F: GTGACGTTGACATCCGTAAGA R: GCCGGACTCATCGTACTCC	NM_007393.5	Mice
Bip	F: ACCACCTACTCGTGC GTTG R: CGTCGAAGACCGTGTCTCA	XM_001927795.7	Swine
Bip	F: GCATCACGCCGTCGTATGT R: ATTCCAAGTACATCCGATGAG	NM_001163434.1	Mice
ATF6	F: ATTCTCCACCTCCCTGTCA R: CCCTGAGTTCCTGCTGATAC	XM_021089515.1	Swine
ATF6	F: CGGTCCACAGACTCGTGTTC R: GCTGTCGCCATATAAGGAAAGG	NM_001081304.1	Mice
CHOP	F: CCCCCTGGAAATGAGGAGGA R: GGAGGTGTGTGTGACCTCTG	NM_001144845.1	Swine
CHOP	F: TATCTTGAGCCTAACACGTCG R: CCAGGTTCTCTCTCCTCAGGT	NM_001290183.1	Mice
ERO1 α	F: TGCTTCTGCCAGGTTAGTGG R: ACAAGGCTTGACAGCACAGT	NM_001137627.1	Swine
ERO1 α	F: TCAAACCCTGCCATTCTGATGA R: CAGAGACTCATCCACGGCTC	NM_015774.3	Mice

Western blot

Radioimmunoprecipitation solution (Solarbio, China) was added to the jejunum and IPEC to accelerate lysis. After centrifuging the samples, the supernatants were collected and mixed with 5× protein loading buffer (Beyotime, China). The target proteins were separated via sodium dodecyl sulfate–polyacrylamide gel electrophoresis and transferred to polyvinylidene difluoride membranes (Biotopped, China). The membranes were sealed with 5% skim milk (Meilunbio, China) at 37 °C for 1 h, and incubated with various antibodies such as anti-β-actin (Bioss, China), anti-immunoglobulin heavy chain binding protein (Bip), anti-activating transcription factor 6 (ATF6), anti-inositol requiring enzyme 1 (IRE1), anti-p-IRE1, anti-protein kinase RNA-like endoplasmic reticulum (ER) kinase (PERK), anti-p-PERK, anti-eukaryotic initiation factor2α (eif2α), anti-p-eif2α, anti-C/EBP homologous protein (CHOP), anti-ER oxidoreductase 1 alpha (ERO1α), anti-pro-cysteine aspartate specific protease 3 (Caspase3), anti-cleaved-Caspase3, anti-B-cell lymphoma 2 (Bcl-2), and anti-Bcl-2 associated X protein (Bax) (Wanleibio, China) at 4 °C overnight. Subsequently, the membranes were incubated with goat anti-rabbit IgG/HRP (Bioss, China) at room temperature for 2 h. The target bands were exposed via supersensitive enhanced chemiluminescence (ECL) reagent (Meilunbio, China) and quantified by an imager (Jiapeng Technology, China). The membranes were washed thrice using PBST as soon as the reagents changed.

Immunohistochemistry

The jejunum tissues of the mice were made into paraffin slices. After dewaxing, the slices were put into citric acid antigen repair solution. H₂O₂ (3%) was added to the slices, which were subsequently incubated at room temperature for 15 min. Then, the slices were blocked with 3% bovine serum albumin (Meilunbio, China) at room temperature for 30 min, incubated with anti-Bip at 4 °C overnight, and enzyme-labelled goat anti-rabbit IgG polymer for 1 h at room temperature. Afterward, the slices were stained with 3,3'-diaminobenzidine and hematoxylin (Origeng, China) for 10 min. Finally, the slices were observed under a fluorescence microscope after dehydration and sealing. The slices were washed three times with PBS or PBST as soon as the reagents changed.

Immunofluorescence

The treated IPEC was fixed in 4% paraformaldehyde (Saiguobio, China) for 30 min, soaked in 0.25% Triton X-100 (Biotopped, China) for 10 min, and sealed with 2% bovine serum albumin for 30 min at room temperature, respectively. Then, the cells were incubated with anti-Bip

at 4 °C overnight, goat anti-rabbit IgG HL/FITC (Bioss, China) at 37 °C for 1 h in the dark, and DAPI (Bioss, China) at room temperature for 5 min. Finally, the cells were observed by fluorescence microscopy. The cells were washed three times with PBS as soon as the reagents were changed.

Statistical analysis

The data were statistically and visually analysed by GraphPad Prism 8.0 and are presented as the means ± SD. These data represent three independent experiments, and the average values were compared by one-way analysis of variance (ANOVA). Post hoc ANOVA was performed using the Duncan method in SPSS version 26.0. *P* < 0.05 was considered to indicate statistical significance. ImageJ 1.46r was used to quantitatively analyse the Western blot bands and the IF fluorescence intensity. The magnification of all fluorescence images was 4× PL FL, and the scale was 1000 μm. IHC images were done at a magnification of 200×.

Results

Purification and endotoxin removal of TsAdSPI

TsAdSPI was expressed and purified as a 44 kDa band, and the endotoxin was removed to eliminate its cytotoxicity (Figure 1).

The optimum dose of H₂O₂ and EN460 for interaction with IPEC

The optimum interaction dose between H₂O₂, EN460, and IPEC was detected using a CCK-8 cell activity detection kit. The results demonstrated that the optimum concentrations of H₂O₂ and EN460 were 100 μM and 16 mM, respectively (Figure 2).

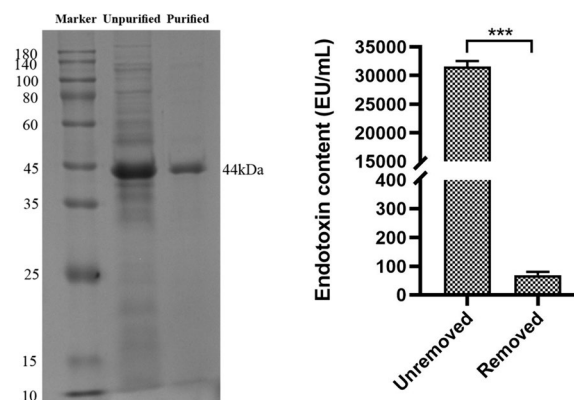


Figure 1 Purification and endotoxin removal of TsAdSPI.

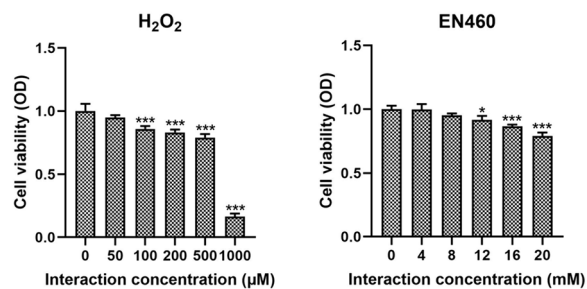


Figure 2 The optimum doses of H₂O₂ and EN460 for interaction with IPEC.

TsAdSPI induced ERS in IECs

To study the effect of TsAdSPI on intestinal ERS, in vivo and in vitro experiments were conducted, and the mice and cells were divided into 10 groups, including the control group, PBS group, 4-PBA group, Tm group, TsAdSPI group, and TsAdSPI+4-PBA group. Changes in ERS-related indexes in the jejunum and IPEC were detected by qPCR, western blotting, IHC, and IF. qPCR revealed that the expression levels of the ERS-promoting genes Bip, ATF6, CHOP, and ERO1 α in the control group and PBS group were not significantly different from those in the 4-PBA group (Figure 3A). Compared with those in the above 3 groups, the transcription levels of ERS-related indexes in the Tm group and TsAdSPI group were significantly greater ($P < 0.001$, $P < 0.01$). The expression of ERS-related indexes in the TsAdSPI+4-PBA group was inhibited in contrast with that in the TsAdSPI group. The western blot results showed that the expression of the ERS-promoting proteins Bip, ATF6, p-IRE1/IRE1, p-PERK/PERK, p-eif2 α /eif2 α , CHOP, and ERO1 α in the TsAdSPI group was significantly greater than that in the control group, PBS group, and 4-PBA group ($P < 0.001$) (Figure 3B). IHC revealed that Bip was mainly expressed in IECs, as shown in the claybank in Figure 3C. The expression in the TsAdSPI group was the highest. The IF results revealed that the expression level of Bip in the TsAdSPI group was significantly greater than that in the PBS group and 4-PBA group ($P < 0.001$) (Figure 3D). These results indicated that TsAdSPI could induce ERS in IECs.

TsAdSPI induced OS in IECs

To investigate the effect of TsAdSPI on intestinal OS, the mice and cells were divided into 10 groups, including the control group, PBS group, NAC group, H₂O₂ group, TsAdSPI group, and TsAdSPI+NAC group. MDA, ROS, and SOD assay kits were used to detect

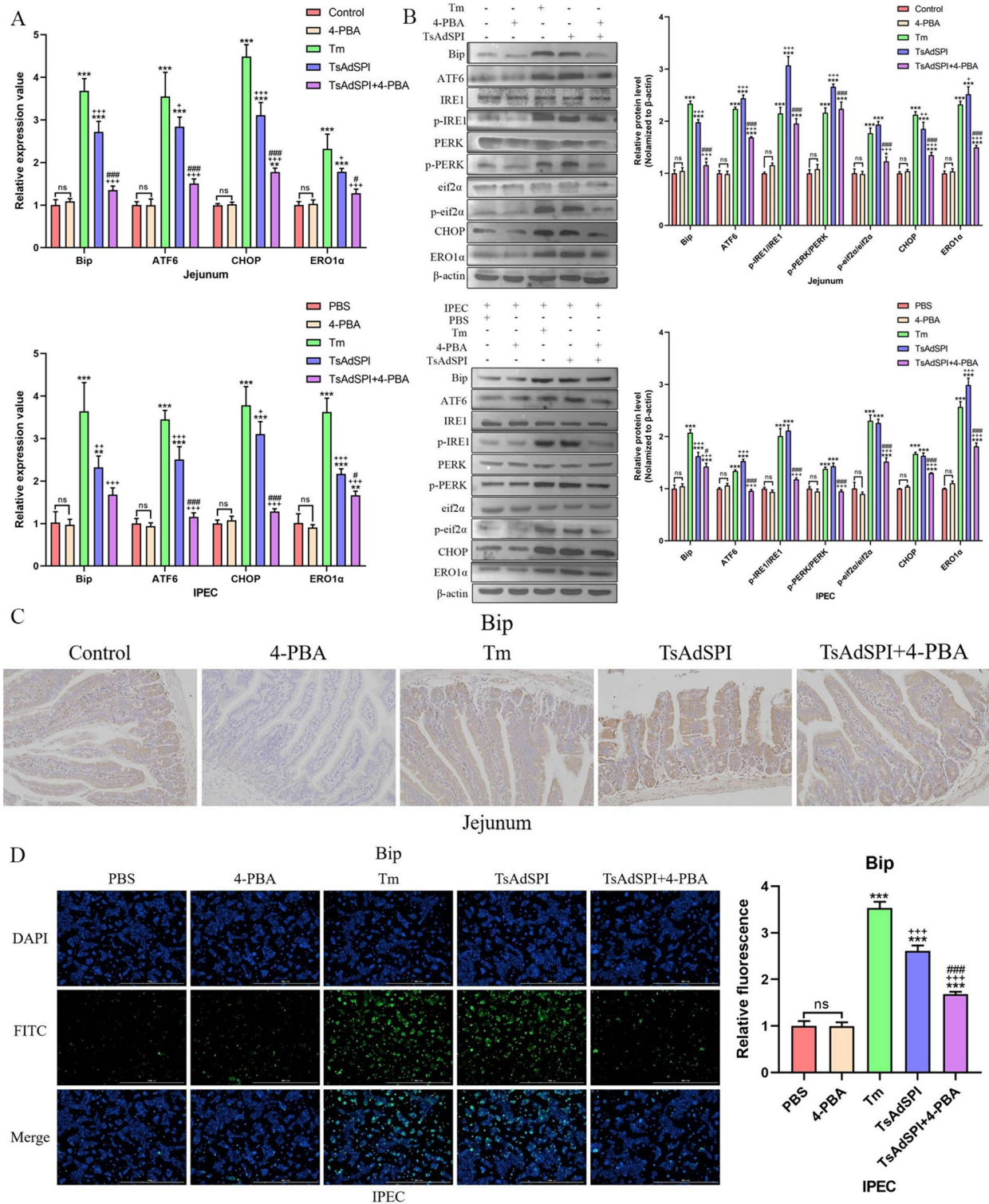
changes in OS-related indexes in the jejunum and IPEC. The results showed that there were no significant differences in OS-related indexes between the NAC group and the control group or PBS group. Figure 4 shows that the content of the oxidation index MDA significantly increased in the H₂O₂ group and TsAdSPI group in contrast with the above 3 groups ($P < 0.001$). The activity of the antioxidant index SOD was opposite to that of MDA. The fluorescence results of the oxidation indexes ROS showed that TsAdSPI caused a significant increase in ROS content ($P < 0.001$). These results indicated that TsAdSPI could induce OS in IECs.

TsAdSPI-induced ERS enhanced OS by activating the PERK-eif2 α -CHOP-ERO1 α axis

To explore whether TsAdSPI-induced ERS affects OS in IECs, the mice and cells were divided into 8 groups, namely, the control group, PBS group, 4-PBA group, TsAdSPI group, and TsAdSPI+4-PBA group, and OS-related indexes were detected in the jejunum and IPEC. The results showed that the content of MDA and ROS in the TsAdSPI group increased significantly and the activity of SOD decreased significantly compared with those in the control group and PBS group ($P < 0.001$) (Figure 5). However, after adding the ERS inhibitor 4-PBA, the content of MDA and ROS in the TsAdSPI+4-PBA group was significantly decreased ($P < 0.001$, $P < 0.01$), and the activity of SOD was considerably increased compared with that in the TsAdSPI group ($P < 0.01$, $P < 0.05$). This finding indicated that TsAdSPI-induced OS depended on ERS and that ERS could increase OS.

To further study the ways in which TsAdSPI-induced ERS increased OS, the key proteins ATF6, IRE1 and PERK of the three ERS signalling pathways were inhibited in IPEC. The results showed that OS-related indexes in the TsAdSPI+AEBSF group and TsAdSPI+MKC3946 group were not significantly different from those in the TsAdSPI group after treatment with the ATF6 inhibitor AEBSF and the IRE1 inhibitor MKC3946 (Figures 6A and B). Only after the addition of the PERK inhibitor GSK2606414 did the OS-related indexes of the TsAdSPI+GSK2606414 group change significantly compared with those of the TsAdSPI group ($P < 0.01$, $P < 0.05$) (Figure 6C). These results suggested that TsAdSPI promoted OS by activating the PERK signalling pathway.

Subsequently, further in-depth exploration was conducted, and the expression levels of the downstream PERK genes and proteins eif2 α , CHOP, and ERO1 α in the IPEC were detected via qPCR and western blotting. The results showed that TsAdSPI considerably increased the transcription levels of CHOP and ERO1 α and the expression levels of p-PERK/PERK, p-eif2 α /eif2 α , CHOP, and ERO1 α in comparison to those in



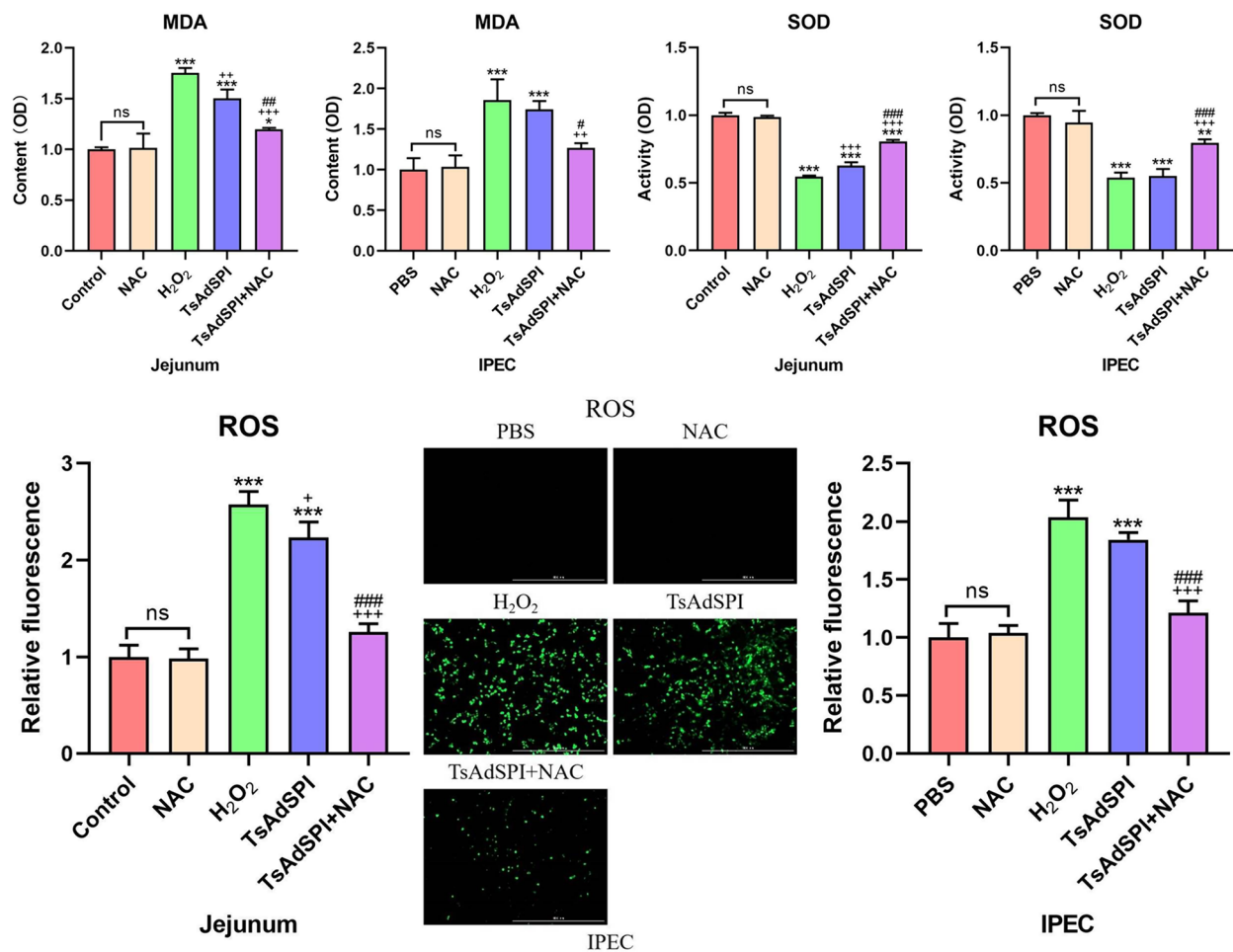


Figure 4 Identification of TsAdSPI-induced OS in IECs. MDA, ROS, and SOD assays were used to detect changes in OS-related indexes. * $P < 0.05$, ** $P < 0.01$, *** $P < 0.001$ compared with the control group, PBS group, and NAC group; + $P < 0.05$, ++ $P < 0.01$, +++ $P < 0.001$ compared with the H₂O₂ group; # $P < 0.05$, ## $P < 0.01$, ### $P < 0.001$ compared with the TsAdSPI group.

the PBS group ($P < 0.001$) (Figure 7A). The expression of these genes and proteins was significantly lower in the TsAdSPI+GSK2606414 group than in the TsAdSPI group after the addition of the PERK inhibitor GSK2606414 ($P < 0.001$). After the addition of the ERO1 α inhibitor EN460, the expression level of ERO1 α in the TsAdSPI+EN460 group decreased significantly ($P < 0.001$), and the expression of ERO1 α 's upstream indexes did not change significantly compared with those in the TsAdSPI group (Figure 7B). Consequently, we speculated that TsAdSPI-induced ERS might enhance OS by activating the PERK-eif2 α -CHOP-ERO1 α axis. The detection results of OS-related indexes confirmed the above conjecture. The contents of MAD and ROS were significantly reduced in the TsAdSPI+EN460 group, while the activity of SOD was significantly upregulated compared with that in the TsAdSPI group ($P < 0.001$, $P < 0.01$).

TsAdSPI-induced OS enhanced ERS by boosting Ca²⁺ efflux
The above results showed that TsAdSPI-induced OS was partially dependent on ERS, but it remains unclear whether TsAdSPI-induced OS is also dependent on ERS. Therefore, the mice and cells were divided into 8 groups, namely, the control group, PBS group, NAC group, TsAdSPI group, and TsAdSPI+NAC group, and the expression of ERS-related indexes was measured via qPCR and western blotting. qPCR results showed that TsAdSPI significantly increased the transcription levels of Bip, ATF6, CHOP, and ERO1 α compared with those in the control group and PBS group ($P < 0.001$) (Figure 8A). Nevertheless, the transcription of these genes was inhibited in the TsAdSPI+NAC group compared with the TsAdSPI group after the addition of the OS inhibitor NAC ($P < 0.001$, $P < 0.01$). The Western blot results displayed a trend similar to that of the qPCR results. The relative expression levels of Bip, ATF6,

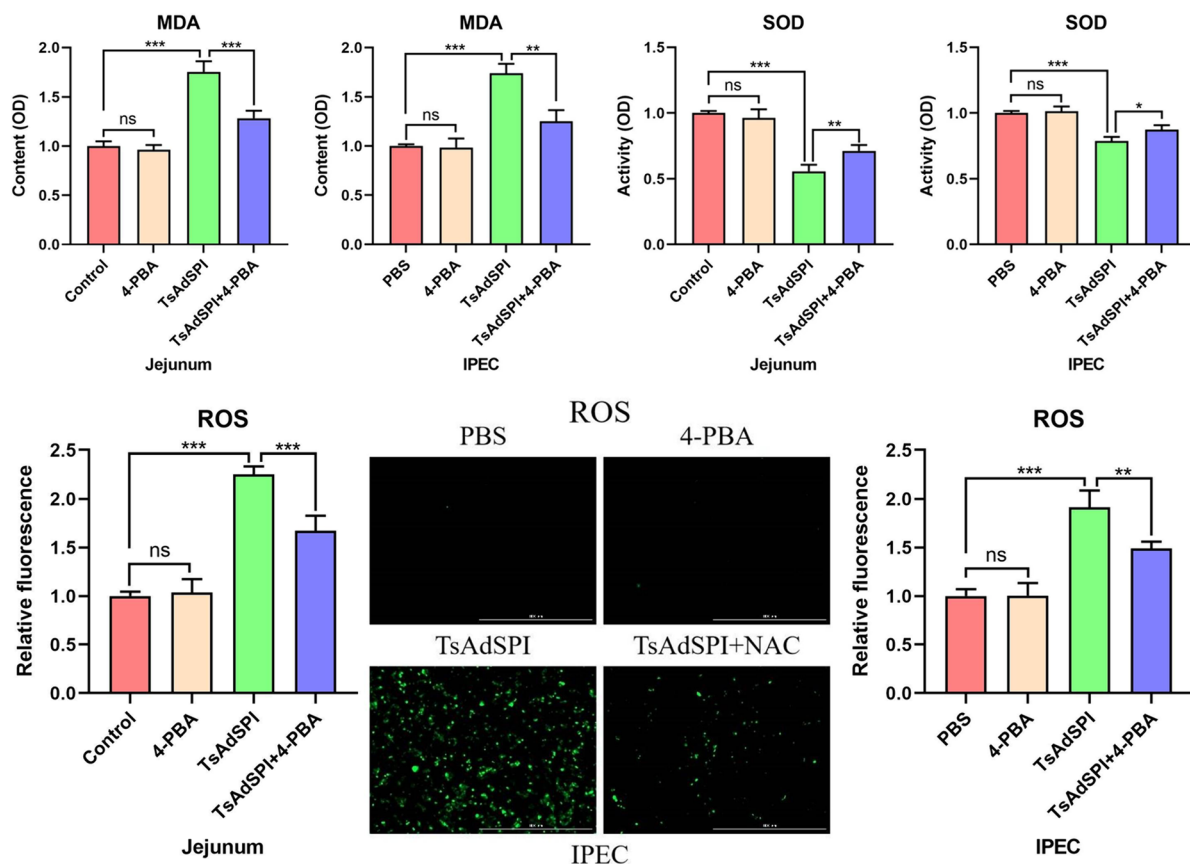


Figure 5 Identification that TsAdSPI-induced ERS enhanced OS.

p-IRE1/IRE1, p-PERK/PERK, p-eif2 α /ef2 α , CHOP, and ERO1 α in the TsAdSPI + NAC group were significantly lower than those in the TsAdSPI group ($P < 0.001$) (Figure 8B). These results demonstrated that the increase in the TsAdSPI-induced ERS level depended on the increase in the OS level, and there was positive feedback between them.

To further study how TsAdSPI-induced OS promoted ERS, the Ca²⁺ level in the IPEC was detected using the Fluo-4 AM fluorescent probe. Figure 9A shows that the concentration of intracellular Ca²⁺ was significantly greater in the TsAdSPI group than in the PBS group ($P < 0.001$). However, the Ca²⁺ concentration in the TsAdSPI + NAC group was significantly lower than that in the TsAdSPI group, indicating that the increase in Ca²⁺ level induced by TsAdSPI was dependent on OS. Subsequently, the Ca²⁺ release antagonist 2-APB was used to further explore the effects of TsAdSPI-induced OS on Ca²⁺ and ERS in IPEC in depth. The qPCR and Western blot results showed that the expression levels of ERS-related genes and proteins in the TsAdSPI + 2-APB group were lower than those in the TsAdSPI group after treatment with 2-APB ($P < 0.001$, $P < 0.01$) (Figures 9B and C).

Accordingly, we hypothesized that TsAdSPI-induced OS increases ERS by inducing Ca²⁺ efflux from the ER.

TsAdSPI-induced OS preceded ERS

To probe the sequence of ERS and OS induced by TsAdSPI, qPCR, western blot, and detection kits were used to observe the changes in ERS- and OS-related indexes 1, 2, 3, 6, and 9 h after IPEC and TsAdSPI were cocultured or not cocultured. Figure 10A shows that there was no significant change in ERS-related indexes after TsAdSPI was used to stimulate IPEC cells for 1 h, but these indexes markedly changed from 2 to 9 h ($P < 0.001$) and peaked at 3 or 6 h. The content of MDA and ROS tended to increase from 1 to 3 h, peaked at 3 h, and then decreased from 3 to 9 h (Figure 10B). The trend of SOD was opposite to that of MDA and ROS and reached its lowest level at 3 h. These results suggested that TsAdSPI-induced OS preceded ERS.

TsAdSPI promoted IEC apoptosis by enhancing ERS and OS

To study the effects of the ERS and OS induced by TsAdSPI on IEC apoptosis, 4-PBA and NAC were used, and the changes in apoptosis-related indexes in the jejunum

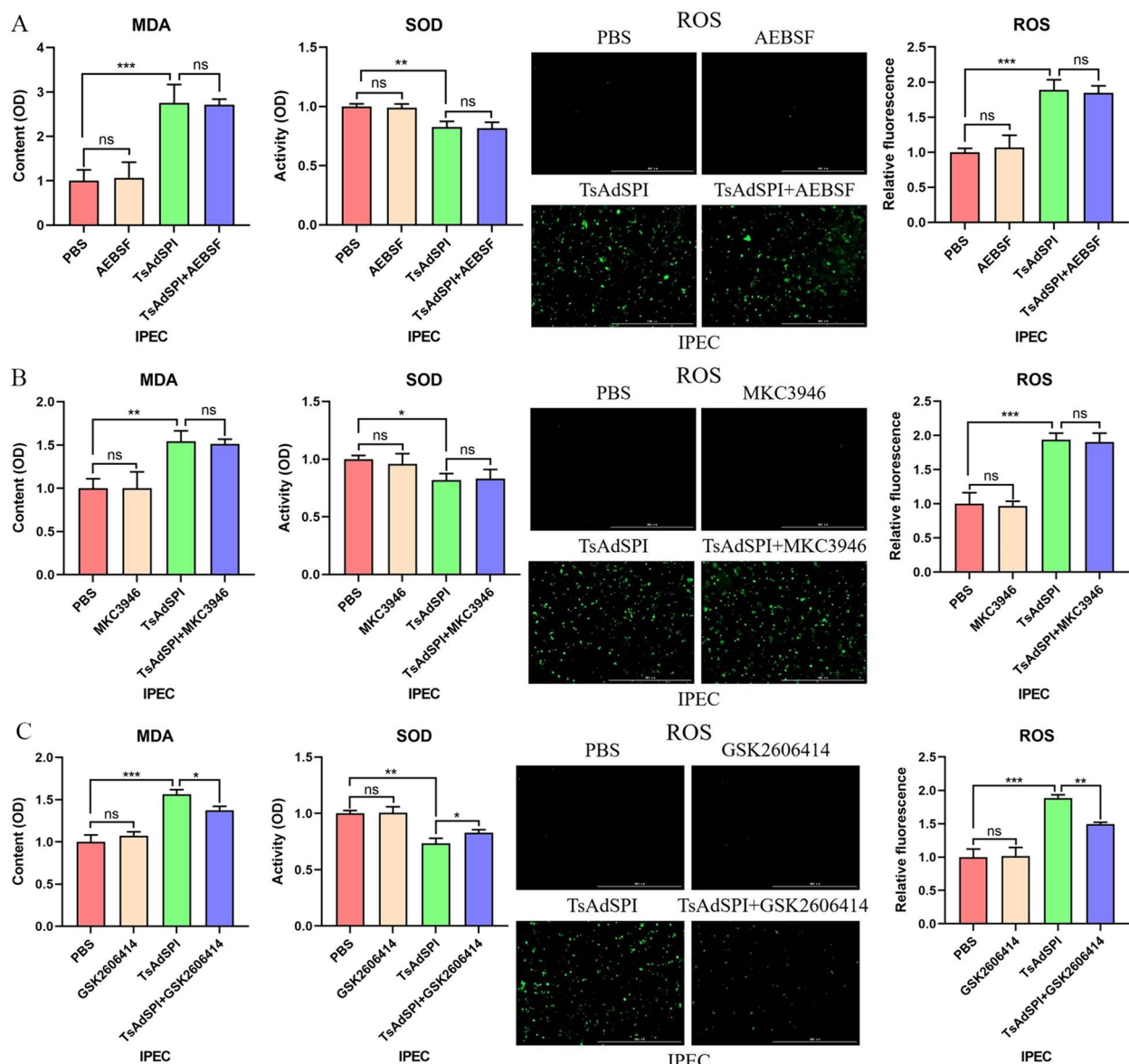


Figure 6 TsAdSPI-induced ERS enhanced OS by activating the PERK signalling pathway. Detection kits for MDA, ROS, and SOD were used to analyse the changes in OS-related indexes after inhibiting ATF6 (A), IRE1 (B), and PERK (C).

and IPEC were measured by western blotting and FCM. As shown in Figure 11A, the expression of the apoptosis-promoting proteins cleaved caspase 3/pro-caspase 3 and Bax in the TsAdSPI group was significantly upregulated ($P < 0.001$), and the expression of the apoptosis-inhibiting protein Bcl-2 was dramatically downregulated compared to that in the control group and PBS group ($P < 0.001$). After the addition of 4-PBA, the expression levels of cleaved caspase-3/pro-caspase-3 and Bax in the TsAdSPI+4-PBA group were significantly decreased, and the expression level of Bcl-2 was significantly increased compared with that in the TsAdSPI group ($P < 0.001$,

$P < 0.05$). Furthermore, the rate of apoptosis in the TsAdSPI+4-PBA group was significantly lower than that in the TsAdSPI group, as shown in Figure 11B ($P < 0.001$). These results showed that TsAdSPI-induced ERS could increase IEC apoptosis. Moreover, the apoptosis-related indexes and the apoptosis rate of the TsAdSPI+NAC group changed significantly compared with those of the TsAdSPI group post-addition of NAC ($P < 0.001$), indicating that TsAdSPI-induced OS could enhance IEC apoptosis (Figure 12).

To further probe the effects of ERS and OS induced by TsAdSPI on early cell apoptosis in IECs, the

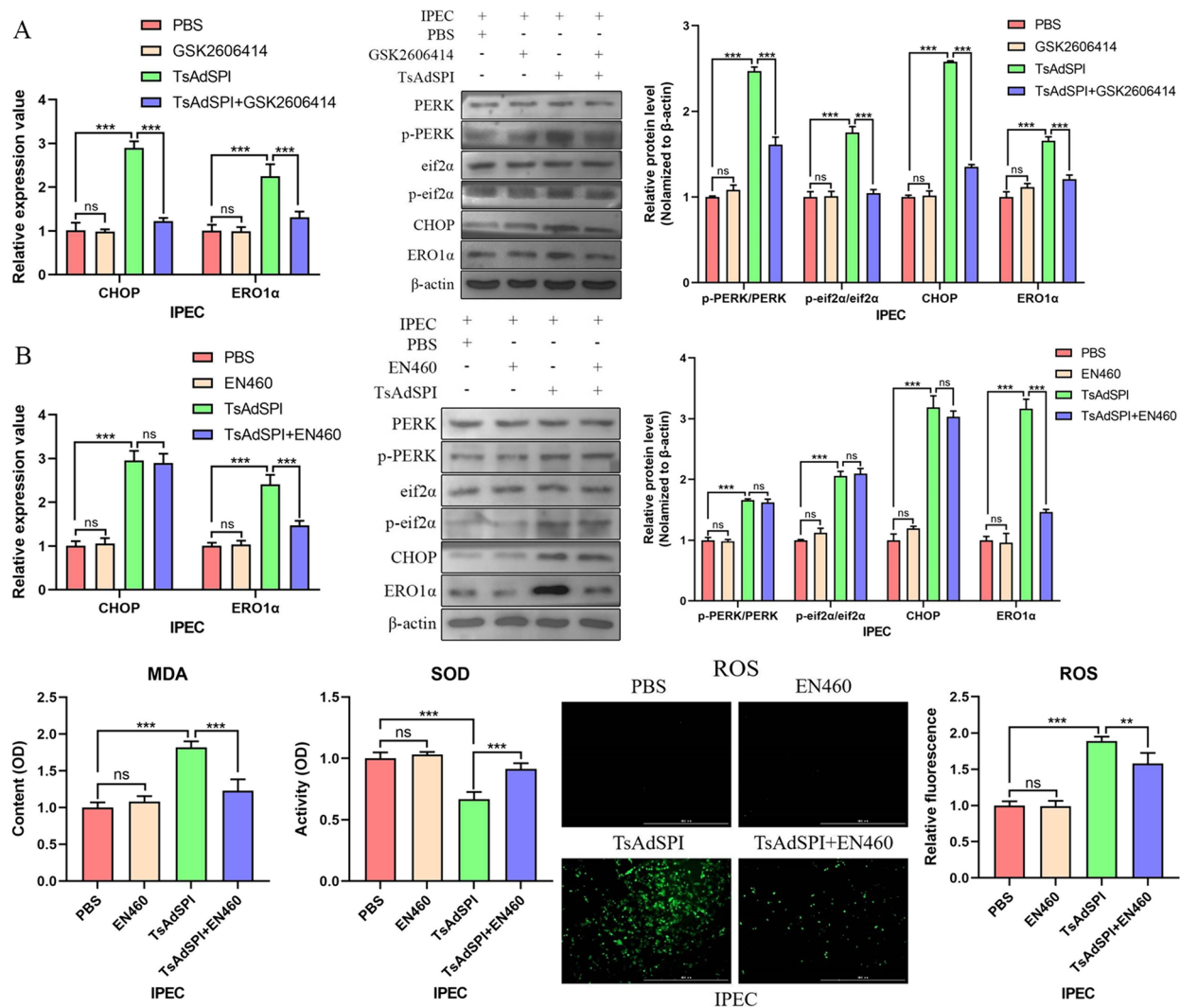


Figure 7 TsAdSPI-induced ERS enhanced OS by activating the PERK-eif2α-CHOP-ERO1α axis. qPCR and Western blot analyses of the expression levels of ERS-related indexes after inhibiting PERK (A). qPCR, Western blot, and detection kits were used to analyse the changes in ERS- and OS-related indexes after inhibiting ERO1α (B).

mitochondrial membrane potential of IPEC cells was detected using a mitochondrial membrane potential detection kit. Figure 13 shows that TsAdSPI decreased the mitochondrial membrane potential, and the decrease in the mitochondrial membrane potential was alleviated by the addition of 4-PBA and NAC. The results revealed that TsAdSPI promoted early IEC apoptosis by enhancing ERS and OS.

Discussion

Many studies have shown that the host reacts to defend against the invasion of exogenous pathogens such as parasites, bacteria, and viruses. However, pathogens can benefit their own survival by inducing ERS and OS in

host cells and achieve a balance between damaging host cells and reproduction [14, 15]. Therefore, on the basis that *T. spiralis* can induce ERS and OS in the host, this study further explored whether *T. spiralis*-secreted TsAdSPI participates in inducing ERS and OS and how to regulate the relationship between them.

ERS is a protective stress process in which cells respond to misfolding, unfolded protein aggregation, and calcium imbalance in the ER cavity and restore ER homeostasis by activating signalling pathways such as the UPR, the ER overload reaction, and apoptosis [10]. As a receptor and molecular chaperone of ER homeostasis, Bip plays an important role in monitoring UPR aggregation and ER activation. During ER homeostasis, Bip binds to the ER

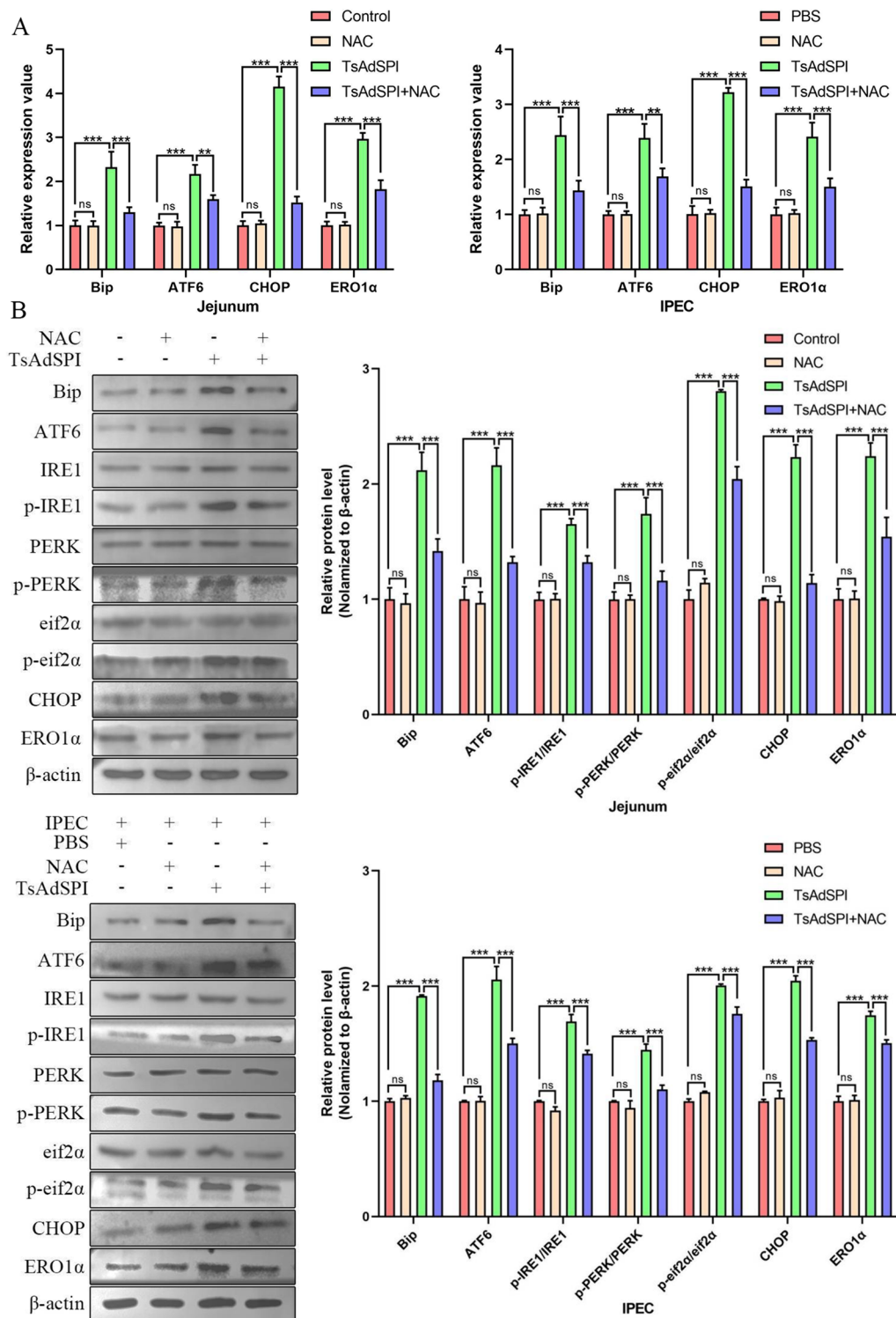


Figure 8 TsAdSPI-induced OS enhanced ERS. qPCR (A) and Western blot (B) analyses of the expression levels of ERS-related indexes.

transmembrane proteins ATF6, IRE1, and PERK. When the UPR increases, Bip has a greater affinity for the UPR and dissociates from ATF6, IRE1, and PERK, activating

three main signalling pathways of the UPR [16]. In our study, Bip, ATF6, IRE1, and PERK and their downstream proteins eif2 α , CHOP, and ERO1 α were detected to

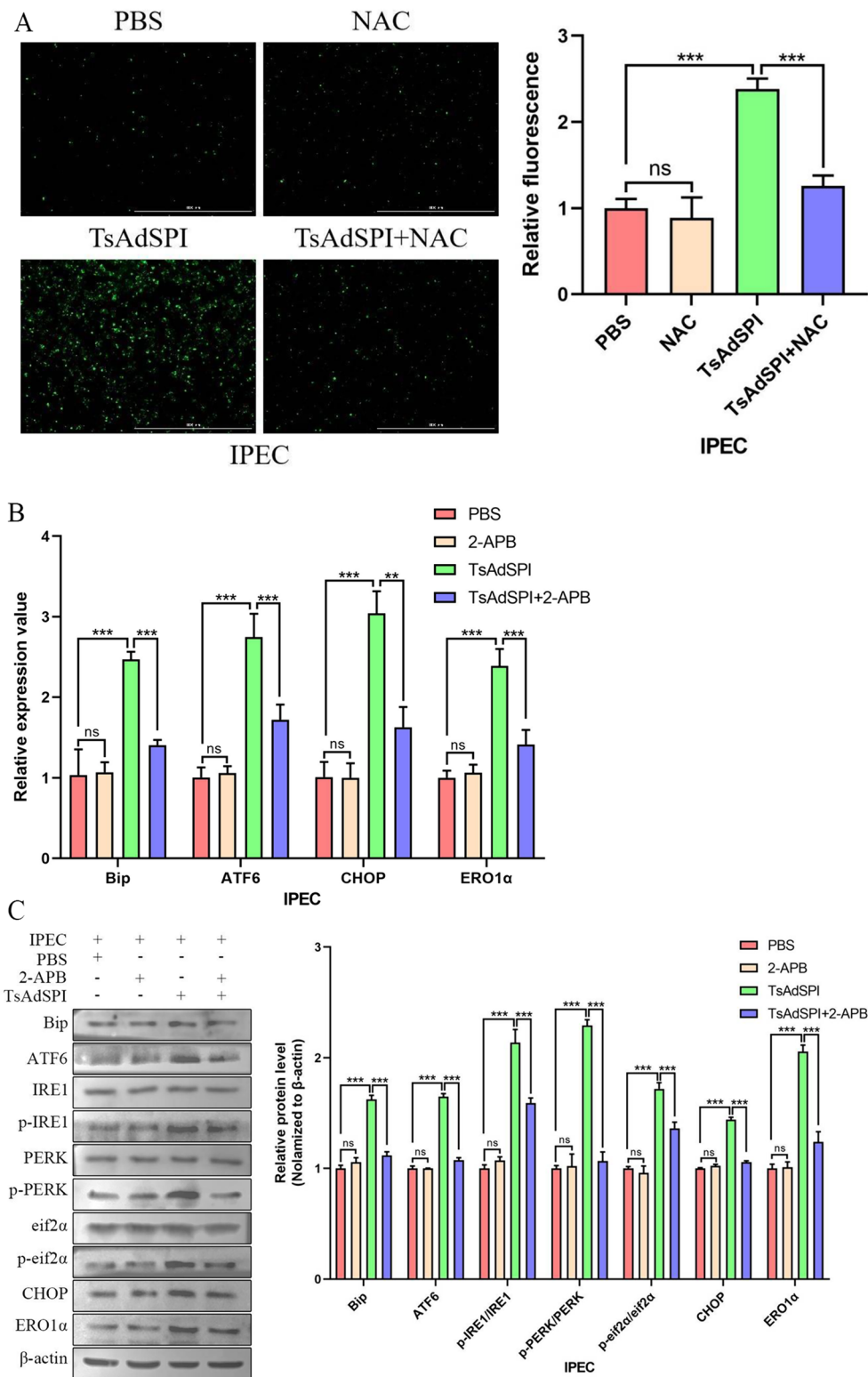


Figure 9 TsAdSPI-induced OS enhanced ERS by boosting Ca^{2+} efflux. The fluo-4 AM fluorescent probe (A) was used to analyse the Ca^{2+} level in IPEC cells, and qPCR (B) and Western blot (C) were used to analyse the expression levels of ERS-related indexes.

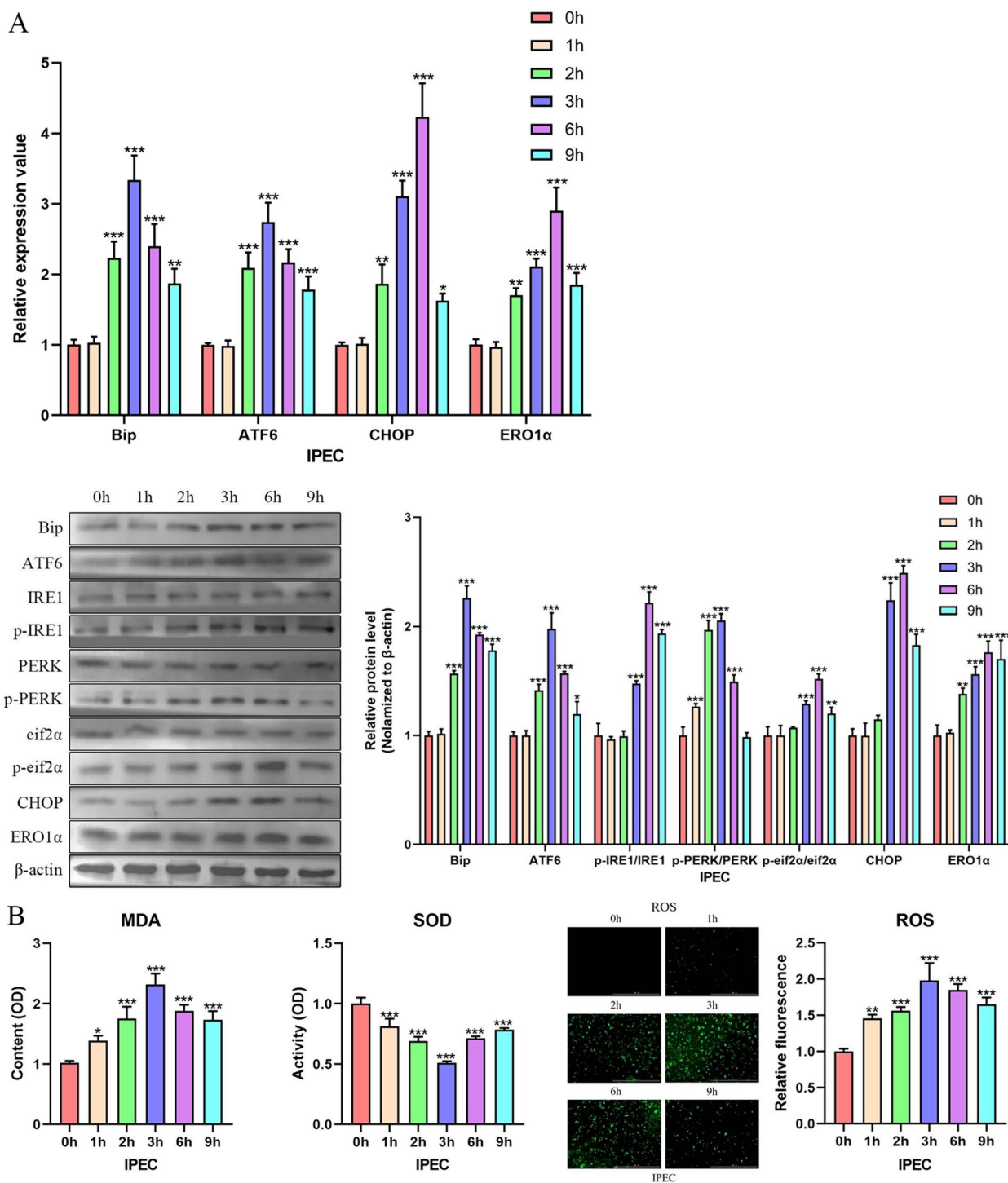


Figure 10 TsAdSPI-induced OS preceded ERS. qPCR and Western blotting (A) were used to analyse the expression levels of ERS-related indexes. The changes in OS-related indexes were analysed with MDA, ROS, and SOD detection kits (B).

determine the effect of TsAdSPI on ERS in the jejunum and IPEC. The results of qPCR, Western blot, IHC, and IF showed that these indexes were significantly upregulated after TsAdSPI stimulation, and Bip was located in

IECs, which indicated that TsAdSPI could induce ERS in host IECs.

When people or animals are stimulated by stressors, the redox system of the body is imbalanced, resulting in

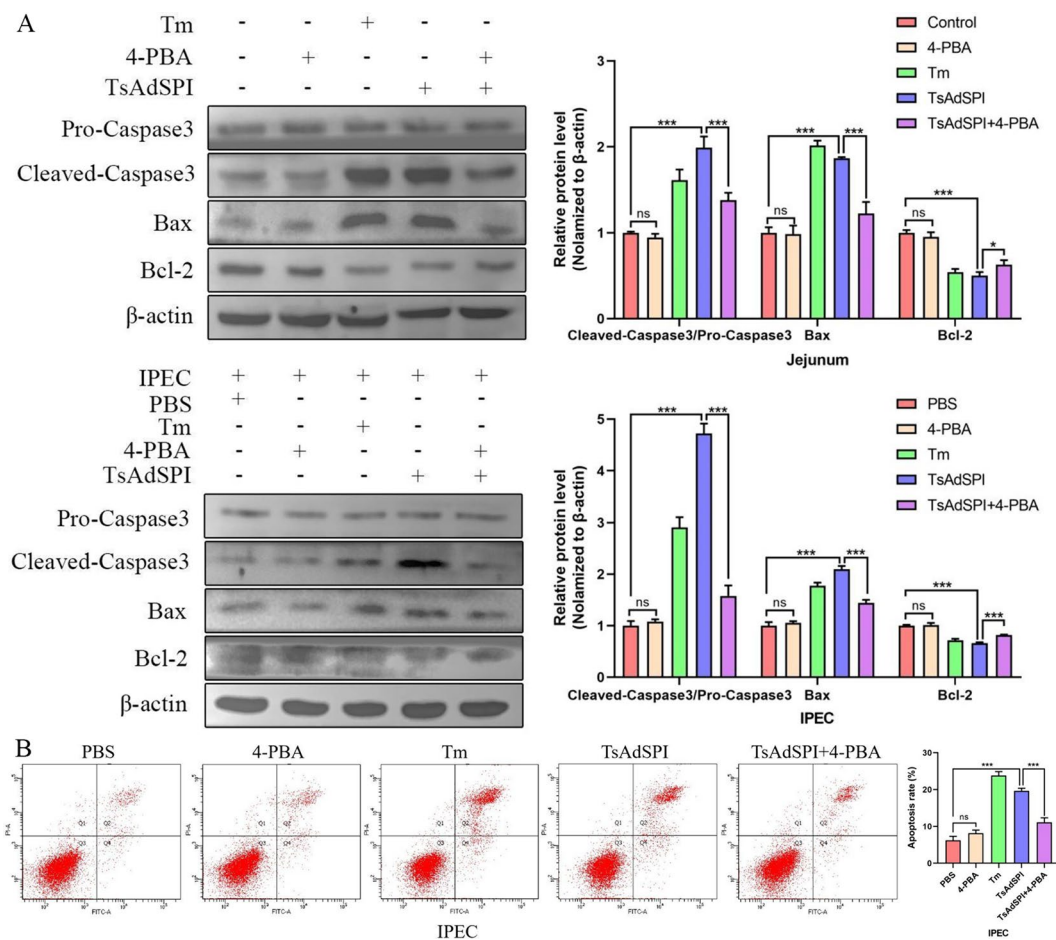


Figure 11 Identification of TsAdSPI promoted IEC apoptosis by enhancing ERS. Western blot (A) and FCM (B) were used to analyse the expression levels of apoptosis-related indexes and changes in the percentage of apoptotic cells.

ROS accumulation and triggering OS. An appropriate amount of ROS can help the body eliminate redundant or harmful cells in a timely manner, which is the normal defense mechanism of the body. Nevertheless, when excessive ROS accumulate, cell homeostasis disorders occur, leading to irreversible cell damage and apoptosis [17]. An increase in the oxidation indexes ROS and MDA and a decrease in the antioxidant index SOD are signs of OS occurrence [18]. Through in vivo and in vitro experiments, we found that ROS, MDA, and SOD in IECs exhibited the same changes as those described above after TsAdSPI stimulation, indicating that TsAdSPI can induce OS in IECs.

Previous results have shown that TsAdSPI can enhance ERS and OS in IECs; therefore, whether there is a connection between them and whether TsAdSPI-induced ERS can affect OS remain unclear. To clarify this, an ERS inhibitor was used, and OS-related indexes were detected. When ERS was inhibited, OS was also inhibited, which indicated that TsAdSPI-induced OS

depended on ERS and that ERS could enhance OS. Moreover, the key proteins ATF6, IRE1, and PERK in the three ERS pathways were inhibited, and OS was only regulated by PERK, while ATF6 and IRE1 had no regulatory effect on OS. We subsequently inhibited PERK and detected the expression levels of the downstream PERK genes and proteins eif2 α , CHOP, and ERO1 α . The results showed that the expression of these three indexes was decreased, and the levels of ROS and MDA were also decreased. This indicated that TsAdSPI-induced ERS could promote OS by activating the PERK-eif2 α -CHOP-ERO1 α axis. Zhou et al. [19] found that *Porcine epidemic diarrhea virus* disrupted the redox homeostasis of the host by manipulating the PERK-CHOP-ERO1 α -ROS axis, which was beneficial for self-replication. This confirmed our results to some extent.

Furthermore, we explored the effect of TsAdSPI-induced OS on ERS. The results showed that the expression of ERS-related indexes decreased significantly after treatment with the OS inhibitor, suggesting that

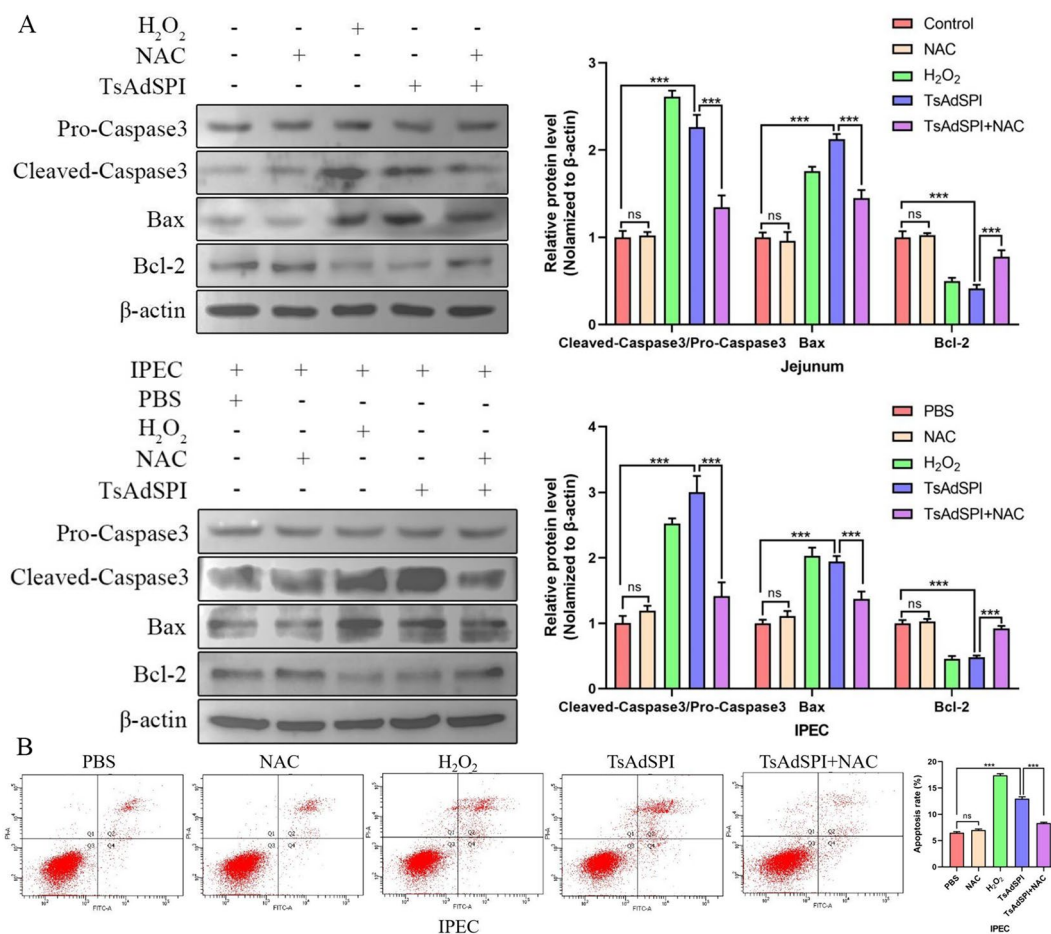


Figure 12 TsAdSPI promoted IEC apoptosis by enhancing OS. Western blot (A) and FCM (B) analyses of the expression levels of apoptosis-related indexes and changes in the percentage of apoptotic cells.

TsAdSPI-induced OS depended on ERS and that there was a synergistic relationship between them. Zeeshan et al. [20] reported that calcium is a redox signalling medium that regulates the relationship between ERS and OS. For this reason, we measured the concentration of Ca²⁺ in the IPEC to determine whether TsAdSPI-induced OS boosted ERS by affecting calcium transfer. The results showed that the intracellular Ca²⁺ level decreased after OS was inhibited. After the addition of a Ca²⁺ releasing antagonist, the expression of ERS-related indexes decreased significantly. This revealed that TsAdSPI-induced OS could strengthen ERS by promoting Ca²⁺ efflux.

Subsequently, the sequence of ERS and OS was determined by detecting the changes in ERS- and OS-related indexes after TsAdSPI was used to stimulate IPEC for different durations. The results showed that OS-related indexes changed significantly at 1 h and peaked at 3 h post stimulation, but the expression of ERS-related indexes reached their highest levels at 3 or 6 h, indicating

that TsAdSPI-induced OS preceded ERS. Thus, we speculated that the IECs produced OS after TsAdSPI stimulation, which caused a redox imbalance in the ER, resulting in the disturbance of ER function and triggering ERS. Moreover, the occurrence of ERS produces a large amount of ROS and exacerbates OS. ERS and OS showed a relationship of mutual correlation and promotion.

Apoptosis is activated when cells are under continuous or excessive stress. This adaptive regulatory process can be activated by signalling pathways such as the IRE1-Caspase12, PERK-Eif2α-CHOP, ROS-fatty acid synthase ligand-Caspase8, and ROS-apoptotic signal regulated kinase 1 pathways [21, 22]. Some studies have shown that the ERS and OS mediated by parasites can induce the apoptosis of host cells [12, 23]. Hence, we speculated that TsAdSPI was involved in the apoptosis of host cells induced by *T. spiralis*-mediated ERS and OS. The experimental results confirmed the above conjecture: TsAdSPI-induced apoptosis was inhibited after inhibiting ERS and OS. This indicated that

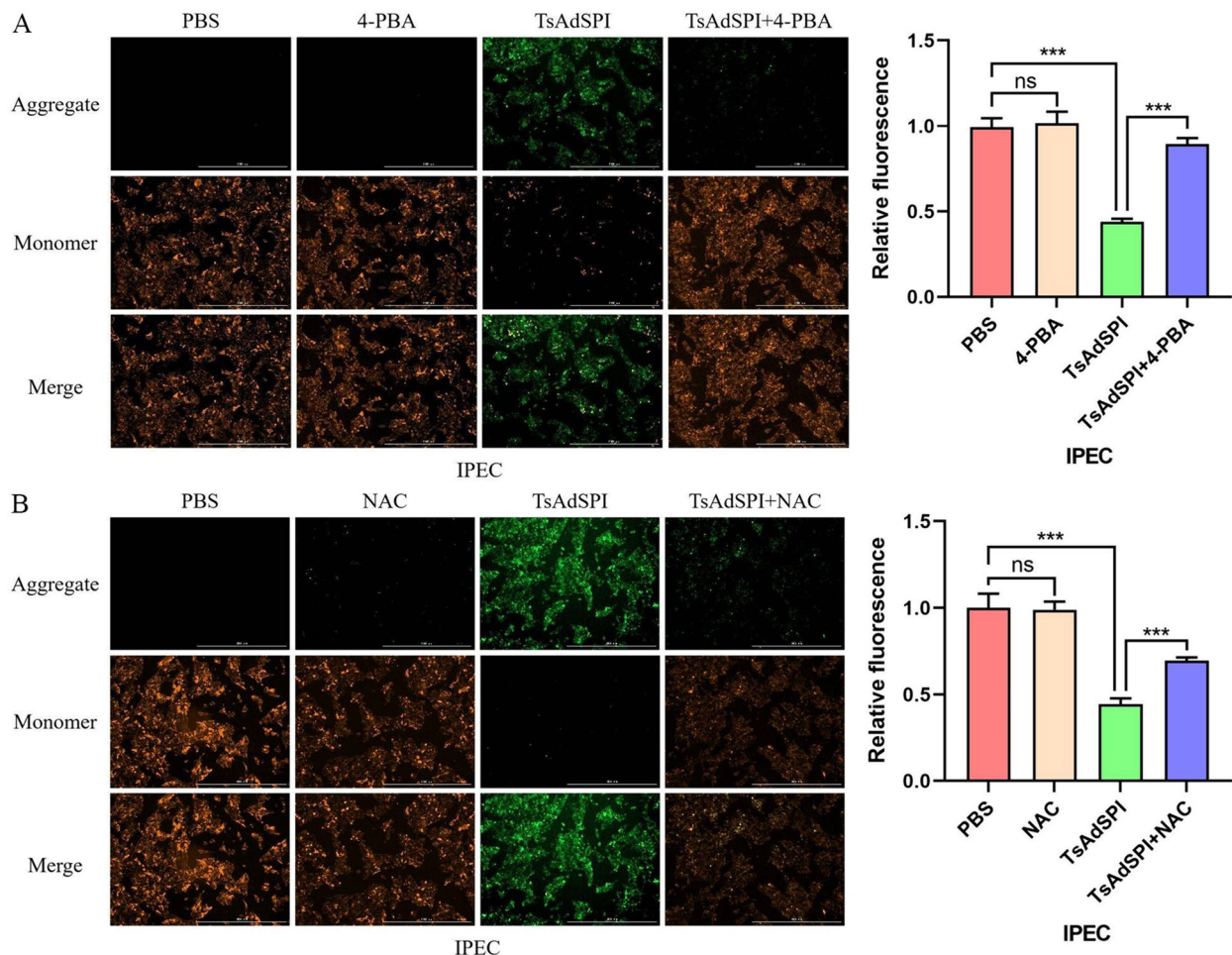


Figure 13 TAdSPI promoted early IEC apoptosis by enhancing ERS and OS. A mitochondrial membrane potential detection kit was used to analyse the mitochondrial membrane potential of IPEC cells after inhibiting ERS (A) and OS (B).

TsAdSPI-induced IEC apoptosis depended on ERS and OS and that ERS and OS were closely related to apoptosis.

In conclusion, after TsAdSPI induced OS after entering IECs, OS promoted ERS by enhancing Ca^{2+} efflux from the ER, and ERS subsequently strengthened OS by activating the PERK-eif2 α -CHOP-ERO1 α axis. The combination of ERS and OS induced by TsAdSPI synergistically promoted IEC apoptosis. These processes may help *T. spiralis* achieve a balance between host cell damage and its own survival and contribute to its parasitism.

Acknowledgements

We thank Northeast Agricultural University and the Harbin Veterinary Research Institute.

Authors' contributions

JZ and LL designed this study. JZ, LL, ZL, FS, YH, QL, YY, XL, JY, and QZ performed the experiments. JZ drafted and revised the manuscript. All the authors have read and approved the final manuscript.

Funding

This work was supported by the National Natural Science Foundation of China (31372427).

Availability of data and materials

The datasets used or analysed during the current study are available from the corresponding author upon reasonable request.

Declarations

Ethics approval and consent to participate

The feeding and experimental procedures for the mice were carried out in accordance with the Chinese Animal Management Ordinance. The animal experiment standards were approved by the Animal Management Committee of Northeast Agricultural University and implemented in accordance with animal ethics guidelines and approved protocols (Animal Ethics Committee approval number SYXK [Hei] 2016-007).

Competing interests

The authors declare that they have no competing interests.

Author details

¹Heilongjiang Provincial Key Laboratory of Zoonosis, College of Veterinary Medicine, Northeast Agricultural University, 600 Changjiang Street, Harbin 150030, China.

Received: 12 March 2024 Accepted: 4 May 2024

Published online: 14 June 2024

References

- Zhen JB, Wang RB, Zhang YH, Sun F, Lin LH, Li ZX, Han Y, Lu YX (2023) Effects of *Trichinella spiralis* and its serine protease inhibitors on autophagy of host small intestinal cells. *Infect Immun* 91:e0010323
- Liu YD, Liu JC, Wang N, You XH, Yang YM, Ding J, Liu XL, Liu MY, Li C, Xu N (2024) Quantitative label-free proteomic analysis of excretory-secretory proteins in different developmental stages of *Trichinella spiralis*. *Vet Res* 55:4
- Xu JY, Pang ZX, Zhang JP, Xia S, Wang RB, Zhang YH, Zhen JB, Song XW, Lin LH, Sun F, Xuan XX, Lu YX (2022) Regulatory effects of *Trichinella spiralis* and a serine protease inhibitor on the endoplasmic reticulum stress response of intestinal epithelial cells. *Vet Res* 53:18
- Molehin AJ, Gobert GN, McManus DP (2012) Serine protease inhibitors of parasitic helminths. *Parasitology* 139:681–695
- Xu JY, Yu PC, Wu LJ, Liu MX, Lu YX (2019) Regulatory effect of two *Trichinella spiralis* serine protease inhibitors on the host's immune system. *Sci Rep* 9:17045
- Cervantes-Ortiz SL, Cuervo NZ, Grandvaux N (2016) Respiratory syncytial virus and cellular stress responses: impact on replication and physiopathology. *Viruses* 8:124
- Zhou YS, Zhou XJ, Dong WY, Zhang YX, Du J, Zhou XD, Fang WH, Wang XD, Song HH (2022) Porcine circovirus type 2 induces CHOP-ERO1 α -ROS-mediated apoptosis in PK-15 cells. *Vet Microbiol* 273:109548
- Ríos-Ocampo WA, Navas MC, Faber KN, Daemen T, Moshage H (2019) The cellular stress response in hepatitis C virus infection: A balancing act to promote viral persistence and host cell survival. *Virus Res* 263:1–8
- Dolai S, Adak S (2014) Endoplasmic reticulum stress responses in *Leishmania*. *Mol Biochem Parasitol* 197:1–8
- Peng M, Chen F, Wu ZD, Shen J (2021) Endoplasmic reticulum stress, a target for drug design and drug resistance in parasitosis. *Front Microbiol* 12:670874
- Chandramathi S, Suresh K, Shuba S, Mahmood A, Kuppusamy UR (2010) High levels of oxidative stress in rats infected with *Blastocystis hominis*. *Parasitology* 137:605–611
- Yu YR, Deng MJ, Lu WW, Zhang JS, Jia MZ, Huang J, Qi YF (2014) Endoplasmic reticulum stress-mediated apoptosis is activated in intestines of mice with *Trichinella spiralis* infection. *Exp Parasitol* 145:1–6
- Elgendy DI, Othman AA, Saad MAH, Soliman NA, Mwafy SE (2020) Resveratrol reduces oxidative damage and inflammation in mice infected with *Trichinella spiralis*. *J Helminthol* 94:e140
- Lee J, Song CH (2021) Effect of reactive oxygen species on the endoplasmic reticulum and mitochondria during intracellular pathogen infection of mammalian cells. *Antioxidants* 10:872
- Tardif KD, Waris G, Siddiqui A (2005) Hepatitis C virus, ER stress, and oxidative stress. *Trends Microbiol* 13:159–163
- Cao SS (2015) Endoplasmic reticulum stress and unfolded protein response in inflammatory bowel disease. *Inflamm Bowel Dis* 21:636–644
- Razavi SM, Soltan MS, Abbasian K, Karami A, Nazifi S (2023) Host oxidative stress in piroplasmosis: a review in domestic animals. *Vet Parasitol* 322:110011
- Chung LY, Wang LC, Chen CH, Lin HY, Yen CM (2010) Kinetic change of oxidative stress in cerebrospinal fluid of mice infected with *Angiostrongylus cantonensis*. *Redox Rep* 15:43–48
- Zhou YS, Zhang YX, Dong WY, Gan SQ, Du J, Zhou XD, Wang WH, Wang XD, Song HH (2023) Porcine epidemic diarrhea virus activates PERK-ROS axis to benefit its replication in Vero E6 cells. *Vet Res* 54:9
- Zeeshan HMA, Lee GH, Kim HR, Chae HJ (2016) Endoplasmic reticulum stress and associated ROS. *Int J Mol Sci* 17:327
- Fu XJ, Cui JJ, Meng XJ, Jiang PY, Zheng QL, Zhao WW, Chen XH (2021) Endoplasmic reticulum stress, cell death and tumor: association between endoplasmic reticulum stress and the apoptosis pathway in tumors. *Oncol Rep* 45:801–808
- Sinha K, Das J, Pal PB, Sil PC (2013) Oxidative stress: the mitochondria-dependent and mitochondria-independent pathways of apoptosis. *Arch Toxicol* 87:1157–1180
- Guha M, Kumar S, Choubey V, Maity P, Bandyopadhyay U (2006) Apoptosis in liver during malaria: role of oxidative stress and implication of mitochondrial pathway. *FASEB J* 20:1224

Publisher's Note

Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.