Abstract. The underlying mechanism involved in auditory neuropathy spectrum disorder (ANSd) remains largely unclear. It has been previously reported that mutations in the apoptosis-inducing factor (AIF) gene are associated with auditory neuropathy and delayed peripheral neuropathy, which can eventually cause ANSD. In the present study, the regulatory effects of AIF knockdown on the cellular functions of spiral ganglion neurons (SGNs) and the molecular mechanism(s) of AIF knockdown in inducing cell apoptosis in SGNs were further investigated. The results showed that the AIF knockdown via siRNA transfection resulted in high levels of oxidative stress, and impaired mitochondrial respiration activity and membrane potential in SGNs. Western blotting further proved that the knockdown of AIF can decrease the content of anti-apoptotic and anti-oxidative proteins, as well as mitochondrial respiratory chain Complex I proteins. The present experimental data suggested that the abnormal expression of AIF may affect SGNs cellular function, and may contribute to the progress of ANSD.

Introduction

Auditory neuropathy spectrum disorder (ANSD) is a specific form of hearing loss that can affect people of all ages (1,2). It is normally characterized by preserved outer hair cell function and abnormal neural conduction of the auditory pathway (1). As a result, the inner ear receives sounds successfully, but the cochlea has a problem with sending the signals from the ear to the auditory nerve and brain stem (3). Impaired hearing presents as fluctuations in hearing sensitivity, as well as speech perception performance, which can eventually lead to social isolation, depression, and a reduction in professional capabilities, especially for young children.

It has been reported that disorders of the cochlea and auditory neuropathies, such as injured inner hair cells (IHCs), abnormal communication between the IHCs and the auditory nerve, or the impaired function of the cochlear spiral ganglion neurons (SGNs) or auditory nerve itself, may account for the pathophysiological mechanisms of ANSD (4,5). However, the underlying mechanism involved in this pathology remains largely unclear.

Several studies have indicated that apoptosis-inducing factor mitochondrion-associated 1 (AIFM1) gene expressing AIFM1 protein or AIF plays a vital role in triggering chromatin condensation and DNA fragmentation to induce programmed cell death (apoptosis), with a significant role in the mitochondrial respiratory chain and metabolic redox reactions (6,7). As a result, the abnormal expression of AIFM1 gene expression is linked to multiple clinical phenotypes, such as mitochondrial encephalomyopathy, spondyloepimetaphyseal dysplasia with mental retardation, X-linked recessive Charcot-Marie-Tooth disease-4 and ANSD (8,9). Recently, a high-level association of mutations was reported in the AIF gene in ANSD. Through the use of whole-exome sequencing (WES), it has been shown that variants in the AIFM1 gene were the main cause of familial and sporadic ANSD (10). Immunostaining of the murine inner ear also indicated a ubiquitous expression of the AIFM1 gene in IHCs, cochlear outer hair cells (OHCs), and especially SGNs. Among these cells, SGNs are the first afferent neurons in the auditory pathway, which makes them the focus of research. However, the potential regulatory effects of AIF knockdown on the cellular function of SGNs are rarely reported.

The aim of the present study was to further study the potential role of AIF dysfunctions in impaired cochlear SNGs cellular functions. The regulatory effects of AIF on mitochondrial function and cellular apoptosis were revealed.
Materials and methods

Tissue culture isolation and detection of AIF expression. The rats were handled in accordance with the NIH Office of Animal Care and Use Animal Research Advisory Committee, and were subjected to the lowest possible levels of pain and discomfort. All animal procedures performed on SPF SD rats were approved by the local Institutional Animal Care and Use Committee of the Chinese PLA General Hospital. Spiral ganglia (n=6 for each time point) were isolated from SD rats on postnatal days 1 (1 d), 4 (4 d), 7 (1 w), 28 (4 w), 56 (8 w) and postnatal year 2. The rats were housed in SPF animal center (temperature 20-25°C) under standard acclimatization conditions of 12 h light/dark cycle and with ad libitum access to water and a commercial standard chow. Health and behavior monitoring by visual examination was performed daily.

The isolation procedures were performed as previously described (11-13). Briefly, for neonatal SD rats (up to 10 days of age), the animals were anesthetized via an intravenous injection of 30 mg/kg sodium pentobarbital and then decapitated. For SD rats older than 10 days, the rats were placed in the euthanasia chamber with readily visible and 100% carbon dioxide was introduced, during which the flow rate must displace no more than 30% of the chamber volume/minute. The expected time to unconsciousness is usually within 2 to 3 min. After observation of each rat for signs of both lack of respiration and faded eye color, the rat was removed from the cage for further tissue isolation.

During the tissue isolation, the skulls were opened along the midsagittal (median) plane with a surgical scissor. Temporal bones from both sides were cut off and placed into ice-cold sterile PBS with 2% BSa and 2% glucose under a stereomicroscope. The cochlear capsule was separated by fine forceps to expose the membranous labyrinth. To avoid cross-contamination, the region where the ganglia were in contact one another was discarded.

To detect the mRNA expression levels of AIF, RNA isolated from the cochlear capsule tissue was measured by RT-qPCR. Briefly, total RNA of the micro-dissected samples was isolated by using an Omega Total RNA kit 1 (Omega Bio-Tek, Inc., Norcross, GA, USA), according to the manufacturer’s instructions. cDNA was synthesized using Takara PrimeScript RT reagent kit with gDNA Eraser (Takara Biotechnology Co., Ltd., Dalian, China), according to the manufacturer’s instructions. RT-qPCR analysis of the RNA was performed using a SYBR Premix EX Taq kit (Takara Biotechnology Co., Ltd.) and RNase-free 96-well PCR plates under the following cycling conditions: 40 Cycles of 95°C for 15 sec and 58°C for 30 sec.

The forward and reverse primers (Sangon Biotech Co., Ltd., Shanghai, China) were used for the specific RNAs in RT-qPCR analysis: AIF, Forward: 5'-CGGCGGTGGTGTGA AAAAAAGAA-3' and 5'-GGAGTCTAGCGGAAACGCC-3' reverse (199 bp); β-actin, Forward 5'-AACCTTCCTGGACTG CCTCC-3' and 5'-CTTCTGACCCATTACCCACC-3' reverse (202 bp). β-actin gene expression was used for normalization and three independent experiments were performed for each time points.

Western blotting. The cochlear capsule tissue (n=3) was lysed with Laemml Sample Buffer (Bio-Rad Laboratories, Inc., Hercules, CA, USA). BCA Protein Assay kit (Beyotime Institute of Biotechnology, Shanghai, China) was used to measure the total protein content of the lysates. The lysates samples were further heated at 100°C for 10 min and loaded onto 10% sodium dodecyl sulfate polyacrylamide gels (SDS-PAGE) with equal amounts of total protein. After being electro-transferred to polyvinylidene fluoride (PVDF) membranes (Bio-Rad Laboratories, Inc.), the samples were blocked with 1X TBST with 5% w/v nonfat dry milk at room temperature for 1 h. The membranes were incubated with primary antibodies overnight at 4°C. Rabbit monoclonal anti-AIF (1:1,000; Abcam, Cambridge, MA, USA), and anti-β-actin (1:4,000; Beyotime Institute of Biotechnology) primary antibodies were used. The blots were then exposed to the horseradish peroxidase-labelled secondary anti-rabbit antibody (1:8,000) for 2 h at room temperature. The proteins were visualized using an enhanced chemiluminescence kit (Thermo Fisher Scientific, Inc., Waltham, MA, USA). The bands signals were detected using a gel imaging system (Syngene, Frederick, MD, USA). The intensities were quantified using ImageJ software and the relatively expression levels of certain proteins were calculated via band intensity normalization.

SGNs isolation and culture. SGNs were isolated from the micro-dissected cochlear capsule tissue of newborn rats (1 day) as described above. The tissues in tube were dissociated in 0.05% trypsin, 0.37 mM EDTA, 5 µg/ml each of collagenase type 4 each, papain, and dispase, and 2,800 U/ml DNase in 1 ml PBS. The tubes were rotated at 37°C for 45 min, with gentle trituration using a pipette every 15 min. After 45 min of enzyme digestion, the tissue was finally triturated 5-10 times and centrifuged for 10 min at 4°C at 800 x g. After discarding the dissociation medium, the cells were re-suspended gently in Dulbecco's modified Eagle's medium supplemented with B272 ml/ml, BDNF 10 µg/ml and penicillin 100,000 U/I 1% (all from Merck KGaA, Darmstadt, Germany) and cultured at 37°C in a humidified incubator with 5% CO2. Cell adhesion and spreading were observed after 12 and 48 h of culture (14,15).

Immunofluorescence. NF200 immunofluorescence was used to identify the isolated SGNs (16). Following culture for 48 h, the SGNs were fixed in 4% paraformaldehyde for 20 min and exposed to 0.2% Triton X-100 in PBS for permeabilization. The cells were then washed with PBS and incubated with 5% BSA blocking solution for 20 min. The SGNs were fixed in 4% paraformaldehyde for 20 min and exposed to 0.2% Triton X-100 in PBS for permeabilization. The cells were then washed with PBS and incubated with 5% BSA blocking solution for 20 min. Following incubation with a rabbit anti-NF200 antibody (1:200; Abcam) overnight at 4°C, the cells were further treated with Alexa-488-conjugated goat anti-rabbit (1:200; Sangon Biotech Co., Ltd.) to label the primary antibodies for 30 min at 37°C. The staining results were observed with a fluorescence microscope (Olympus Corporation, Tokyo, Japan).

siRNA knockdown of AIF in SGNs. To verify the role of AIF in SGNs, we firstly performed siRNA knockdown of AIF using siRNA oligonucleotides (oligos) targeting the coding region of AIF. Following culture for 24 h, the SGNs were transfected with siRNA via Amaxa 4D nucleofector (Lonza Group, Ltd., Basel, Switzerland) using P4 Primary Cell 4D-Nucleofector® X kit L (Lonza Group, Ltd.), according to the manufacturer's instructions. Half of the medium was then replaced with fresh
culture medium after 4 h and the cells were incubated in a humidified incubator with 37°C/5% CO₂ until analysis. The siRNA-AIF oligos for AIF targeting were as follows: Sense 5'-CAGAAAGGAGGCUAGUCCUCAAUdTdT-3', and anti-sense 5'-AUUGGAGGAACUGAGCCUCUCUdGdTdT-3', and siRNA-control oligos for control experiments, Sense 5'-GCAUCGAUUUCUCACCUpdTdT-3', and Anti-Sense 5'-AAGGUAGAAUUCUGAUGCdTdT-3' were used. siRNA (15 pmol/sample) was used, and normal siRNA was replaced of with Cy3 labeled siRNA for observation of the transfection efficiency. The AIF protein contents were analyzed by western blotting. The SGNs cultured on tissue culture polystyrene (TCP) without further treatment were termed the TCP group, which was used as the blank comparison group.

**Intracellular reactive oxygen species (ROS) level detection.** To evaluate the regulatory effect of the AIF siRNA knockdown on cellular oxidative stress status, the intracellular ROS was monitored using 2',7'-dichlorofluorescin diacetate (DCFH-DA). Forty-eight hours after siRNA transfection, the SGNs were treated with DCFH-DA (10 μM) at 37°C for 20 min in the dark. Next, flow cytometry (BD FACS Calibur; Becton, Dickinson and Company, Franklin Lakes, NJ, USA) was performed to measure the fluorescence intensity of the treated cells. The fluorescence of the oxidation product of DCFH-DA, DCF, was excited at 488 nm and detected at 535 nm by flow cytometry. Data was acquired and analyzed using the CELL Quest programme PRO software (Becton, Dickinson and Company). The percentage of cells with a high DCF fluorescence of 10⁻⁷⁻¹⁰⁻⁸ units, as measured by flow cytometry, was used to indicate the intracellular (ROS) level.

In the staining fluorescence of SGNs was also observed with a fluorescent microscope (Leica Microsystems GmbH, Wetzlar). Images from 4 random fields in each group were recorded and the fluorescence density was quantified with ImagePro Plus™ software.

The intracellular malondialdehyde (MDA) levels were determined by using a Lipid Peroxidation MDA Assay kit (Beyotime Institute of Biotechnology). To determine the intracellular malondialdehyde (MDA) levels, the SGNs were lysed with cell lysis buffer and the total protein content of the lysates was measured using BCA Protein Assay kit (Beyotime Institute of Biotechnology). The MDA levels of cell lysis with equal protein contents were detected according to the manufacturer's instructions and expressed as μmol/mg protein.

**Mitochondrial membrane potential.** The mitochondrial membrane potential (Δψm) was assessed using MitoProbe™ DiIC1 (5) according to the MitoProbe™ DiIC1 (5) Mitochondrial Membrane Potential Protocol. The cells were analyzed by flow cytometry with a 633 nm excitation using emission filters appropriate for Alexa Fluor 633 dye. The percentages of cells with a high DiIC1 fluorescence (>10³ units) measured by flow cytometry indicated the mitochondrial membrane potential level.

**Determining of protein contents.** The proteins contents of SGNs 24 h after transfection were determined by western blotting. Immunoblotting was performed using the following primary antibodies: Anti-cytochrome complex (Cyt c) (1:2,000), anti-Bcl-2 (1:1,000), anti-Bax (1:1,000), anti-poly(ADP-ribose) polymerase (PARP)-1, anti-superoxide dismutase 1 (SOD)-1 (1:1,000), anti-SOD-2, anti-Caspase-3 (Casp-3) (1:1,000), anti-C I 39 kDa (1:1,000), anti-C I 20 kDa (1:1,000), anti-C I 17 kDa (1:1,000), anti-C III 49 kDa (1:1,000), anti-C III 47 kDa (1:1,000), anti-C IV 20 kDa (1:1,000), anti-PAR (1:500), anti-GAPDH (1:8,000), anti-β-actin (1:10,000), anti-tubulin (1:10,000), anti-Histone H3 (1:1,000) and anti-COX IV (1:1,000), which were bought from Cell Signaling Technology, Inc., Danvers, MA, USA.

For nuclear, mitochondrial and cytosolic protein enrichment, the cells were fractionated into nuclear, mitochondrial and cytosolic fractions using the Focus SubCell kit (Geno Technology, Inc., St. Louis, MO, USA). Briefly, the treated cells were first harvested and washed in ice-cold SubCell Wash Buffer and then re-suspended in ice cold FOCUS™ SubCell Buffer-I. Following mechanic lysis and the addition of SubCell Buffer-II, the lysed cells were centrifuged for 10 min at 700 x g to pellet the nuclei. The nuclear pellet was further enriched by gradient centrifugation and used for further experiments. The cytosolic supernatant with mitochondria was further centrifuged at 12,000 x g for 15 min at 4°C to pellet the mitochondria for protein content detection. Finally, the cytosolic supernatant was removed and placed into a fresh tube. The enriched proteins were analyzed by SDS-PAGE and western blotting. The relative protein expression levels were calculated by band intensity normalization to internal control, and COX IV, Tubulin and histone H3 are used as internal control for the mitochondrial, cytoplasmic and nuclear fractions respectively (18-20).

**Statistical analysis.** Quantitative data are expressed as the means ± standard deviations (mean ± SD). Statistical analysis was performed using one-way ANOVA followed by Tukey’s post-hoc test for multiple comparisons (using SPSS software). A P<0.05 was considered to indicate a statistically significant difference.

**Results**

**AIF expression decreased during the stage of growth and development.** To characterize the potential role of AIF in the auditory pathway, we firstly determined the expression levels of AIF in cochlea spiral tissue from rats at different growth phases. As shown in Fig. 1A and B, RT-qPCR results indicated that the mRNA expression levels of AIF decreased over time during the stage of growth and development. However, the AIF
expression levels in elderly rats were increased to a certain extent. Western blotting (Fig. 1C and D) results of AIF protein contents confirmed a similar trend of the change.

Isolation and identification of SGNs. Fig. 2A illustrates an example of cultured SGNs with no obvious neurite after 12 h of culture in primary growth medium. More neurites emanated from the cell body after 48 h of culture. Immunofluorescence results indicated that the surviving SGNs were mostly stained with primary anti-nF200 antibody (Fig. 2B) (21).

AIF knockdown increases the oxidative stress status of SGNs. To verify the role of AIF in SGNs apoptosis execution, AIF siRNA knockdown was performed using siRNA oligos targeting the coding region of AIF. As shown in Fig. 3A, the siRNAs labeled with Cy3 fluorescent dye were transfected into SGNs at 24 h after treatment. Compared to normal SGNs or a control siRNA oligo, AIF knockdown significantly decreased the AIF contents in SGNs (Fig. 3B and C).

Twenty-four hours after siRNA knockdown, the levels of reactive oxygen species (ROS) were measured by the cell-permeant fluorescent indicator 2',7'-dichlorodihydrofluorescein diacetate (DCFH-DA). Following treatment with DCFH-DA, intracellular ROS levels were evaluated by flow cytometry and a fluorescent microscope (Fig. 4A-C). As compared to normal SGNs in the TCPS group and control siRNA oligo in the siRNA-control group, AIF knockdown in SGNs significantly increased their intracellular ROS levels.

MDA is a mutagenic product of lipid peroxidation, which is closely associated with intracellular oxidative stress levels.

Generally, cellular MDA levels are low in SGNs, but AIF knockdown could significantly increase them (Fig. 4D), which also confirmed the high oxidative stress status in the siRNA-AIF group.
Figure 3. Transfection of SGNs with siRNA to knockdown AIF expression. (A) Observation of the Cy3-labeled siRNAs 24 h after their transfection into SGNs. (B) Western blotting of AIF content in SGNs 24 h after transfection and (C) its quantitative analysis (n=6). **P<0.01 vs. TCPS and siRNA-control. SNG, spiral ganglion neuron; AIF, apoptosis-inducing factor.

Figure 4. AIF knockdown increases oxidative stress in SGNs. (A) Flow cytometry, (B) fluorescence microscope observation of DCFH-DA staining and (C) quantitative analysis of its fluorescence intensity (n=12). (D) MDA detection of lipid peroxidation in SGNs (n=12). **P<0.01 vs. TCPS and siRNA-control. SNG, spiral ganglion neuron; DCFH-DA, 2',7'-dichlorofluorescein diacetate.
AIF knockdown attenuated mitochondrial respiratory functions and ATP production. Mitochondrial dysfunction is widely implicated in cells dysfunction through impaired ATP synthesis. Since mitochondrial complex I is a major source of mitochondrial ROS (22), the specific activity of complex I in SGNs was also analyzed. As shown in Fig. 5A and D, complex I activity and ATP was significantly decreased following AIF knockdown, yet similar levels of mitochondrial complex II/III (Fig. 5B) and mitochondrial complex IV (Fig. 5C) were observed between the three groups. Flow cytometry results (Fig. 5E) of mitochondrial membrane potential revealed that AIF knockdown markedly decreased the mitochondrial membrane potential of SGNs.

AIF knockdown impairs SGNs functions through anti-oxidative and anti-apoptotic bioactivities. To verify whether AIF knockdown had an effect on anti-oxidative and apoptotic-related factors, western blotting was performed. As
shown in Fig. 6A and B, AIF knockdown in SGNs resulted in the decrease of anti-apoptotic factors cytochrome complex (Cyt C) and Bcl-2, and the increase of pro-apoptotic proteins Bax and Caspase-3 (Casp-3). Although similar contents of PARP and SOD-1 were observed between the three groups, anti-oxidative factor SOD-2 in SGNs was decreased after in SGNs following AIF knockdown. Western blotting analysis (Fig. 7A and B) of mitochondrial respiratory complex-related proteins indicated that AIF knockdown decreased the complex I proteins Complex I 39 kDa and Complex I 20 kDa, but not Complex I 17 kDa, Complex III 49 kDa, Complex III 47 kDa and Complex IV 20 kDa.

Of note, although AIF knockdown led to a decrease in the AIF protein content in SGNs, western blotting (Fig. 8A and B) of enriched mitochondrial, nuclear and cytosolic proteins indicated that mitochondrial siRNA-AIF was decreased, while nuclear and cytosolic siRNA-AIF were observed with the increase in AIF protein. Similar levels of PARP-1 expression was observed in the nuclei of the three groups (Fig. 8A and C). However, the cytosolic and nuclear poly(ADP-ribose) (PAR) polymer levels were increased following AIF knockdown (Fig. 8A and D).

Discussion
An improved understanding of the underlying mechanisms of ANSD is essential for the development of new prevention and treatment mechanisms. Previously studies have suggested that
the impaired cellular functions of SGNs in the cochlea play vital roles in the progression of hearing loss, but the potential mechanism involved remains largely unclear. The pathological factors inducing programmed cell death (apoptosis) may be the potential mechanism underlying SGNs dysfunction and cellular injury. The results the present study revealed new regulatory effects of the AIF protein on SGNs dysfunctions in vitro. The knockdown of the AIF expressions with siRNA resulted in high levels of oxidative stress status, impaired mitochondrial respiration activity and membrane potential in SGNs. Western blotting analysis further indicated that AIF knockdown could decrease the expression levels of anti-apoptotic and anti-oxidative executors, as well as mitochondrial respiratory chain Complex I proteins.

AIF is an NAD(P)H oxidase with oxidoreductase activity. AIF was first identified as a flavoprotein and acts as a mitochondrial effector of apoptosis. Studies have indicated that AIF plays various roles in apoptosis, electron transport, and ferredoxin metabolism, etc. Furthermore, the silencing of the AIF gene was found to increase ROS generation in the mitochondrion (15). ROS overproduction induced high level oxidative stress status, which plays pivotal roles in various pathologies, especially the apoptosis of impaired cells. In the present study, we firstly determined the expression levels of AIF in cochlea spiral tissue from rats at the different growth phase to primary characterize the potential role of AIF in auditory pathway. The results indicated the decreased AIF expression levels of rats as it grows older (during the growth and development stage). Thus, we further investigated the potential role of AIF knockdown in cochlear spiral ganglion neurons (SGNs) cellular functions. As a result, AIF knockdown resulted in a significant overproduction of ROS, as proven by DCFH-DA staining, and high levels of lipid peroxidation, as measured by MDA. Western blotting results showed that anti-apoptotic executors Cyt C and Bcl-2, and anti-oxidative executor SOD-2, were decreased in SGNs by AIF knockdown.
while pro-apoptotic protein Bax and Casp-3 were increased. The high oxidative stress status could potentially lead to injury and apoptosis in SGNs, which may result in dysfunctions of SGNs.

Normally, the flavoprotein AIF is contained in the mitochondrial intermembrane space or inner mitochondrial membrane (23). Upon the induction of apoptosis, AIF translocates from the mitochondria to the cytosol, and rapidly relocates to the nuclear compartment, where it can mediate the fragmentation of large-scale DNA (24). Although AIF is not a component of complex I, AIF-deficiency has been proven to reduce complex I content, indicating its important roles in the biogenesis and/or maintenance of mitochondrial respiratory chain complexes function (25). The present results indicated that AIF knockdown decreased the complex I activity, ATP generation and mitochondrial membrane potential in SGNs. These results suggested that AIF knockdown may induce the mitochondrial dysfunction.

Of note, although it remains unknown what it is that can regulate the release of AIF, in the present study, the data suggested that, even though AIF knockdown resulted in a decrease of the total AIF protein in SGNs, the release and translocation of AIF from the mitochondria resulted in high levels of nuclear and cytosolic AIF. As a result, although the contents of PARP-1 in the nuclei were similar among the three groups, the PAR polymer levels, a product of activated PARP-1, were increased in the AIF-knockdown group (26,27). However, it remains unknown how AIF knockdown increase the levels of PAR, which conversely leads to AIF release.

In addition, although the expression levels of AIF in cochlea spiral tissue were found to be decreased over time during the stage of growth and development, they were
shown to be increased, to a certain extent, in the elderly. The variation of AIF expression levels might be closely associated with the complicated regulatory roles of AIF expression at different physiological phases, be that its vital mitochondrial apoptosis inducing factor (AIF) is an FAD-containing and NADH oxidase and critically important for energy metabolism (29,30). Many studies indicated that AIF may regulate mitochondrial function by participating in the assembly and/or stabilization of the respiratory complexes and contributed to the normal cellular functions. However, similar to other mitochondrial factors, the AIF can also be released from mitochondria during apoptosis, and then migrates to the nucleus, inducing chromatin condensation and DNA fragmentation by unknown molecular mechanism. As a result, the AIF, as seen in cytochrome c, may also has a bifunctional protein with dissociable apoptogenic and redox properties (31). Thus, we speculated that the AIF might play certain positive roles during the early growth and development stage. However, over time, the potential positive roles of AIF might gradually reduce, and the expression levels of AIF were decreased by unknown molecular mechanism. The decrease of AIF expression levels might lead to the AIF deficiency. The AIF further compromises oxidative phosphorylation and contributed to the dysfunction and apoptosis of targeted cells. But, as the rats get older, AIF might play much more important role in the apoptosis inducing, and the increase of some apoptotic factors might enhance the expression of AIF, yet with negative control results of AIF. We thought it will be important to investigate different potential roles of AIF during development and aging, in particular, to what extent and through which exact mechanisms contributes to regulation the expression levels of AIF as well as its exactly regulation roles. What's more, the new experiment methods, such as site-specific gene delivery system, new CRISPR/Cas 9 Technology or site injection of AIF-activating or AIF-inhibiting molecules could be used for AIF gene regulation (knockdown or overexpression) study as well as its in vivo potential regulation effects.

In conclusion, the present data revealed new regulatory effects of the AIF protein on SGNs dysfunctions in vitro. It was shown that AIF knockdown with siRNA resulted in high levels of oxidative stress status, impaired mitochondrial respiration activity and membrane potential in SGNs. Western blotting analysis further indicated that AIF knockdown could decrease the expression levels of anti-apoptotic and anti-oxidative executors, as well as mitochondrial respiratory chain Complex I proteins. This study provided a molecular basis for the development of new prevention and therapeutic strategies for ANSD.

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Availability of data and materials

The datasets generated and/or analyzed during the current study are not publicly available due to the institutional rules of the PLA General Hospital but are available from the corresponding author on reasonable request.

Authors’ contributions

LZ and JZ performed the experiments and drafted the manuscript. WW designed the experiments. JW and DH conducted statistical analysis of the data. ML conceived the study, and participated in its design and coordination.

Ethics approval and consent to participate

All animal procedures performed on SPF SD rats were approved by the local Institutional Animal Care and Use Committee of the Chinese PLA General Hospital.

Patient consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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