Down-regulated RPS-30 in Angiostrongylus cantonensis L5 plays a defensive role against damage due to oxidative stress

Wei-Wei Sun  
Wenzhou Medical University

Xiu-Mei Yan  
Wenzhou Medical University Second Affiliated Hospital

Qing Shi  
Wenzhou Medical University

Yuan-Jiao Zhang  
Wenzhou Medical University

Jun-Ting Huang  
Wenzhou Medical University

Hui-Cong Huang  
Wenzhou Medical University

Hong-Fei Shi  
Nanyang Normal University

Bao-Long Yan (✉️ 1982yblblog@163.com)  
Wenzhou Medical University  https://orcid.org/0000-0001-9231-3327

Research

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Abstract

Background: Eosinophilic meningitis, caused by Angiostrongylus cantonensis L5, is mainly attributed to the Eosinophils, which contribute to tissue inflammatory responses in helminthic infections. Eosinophils are associated with helminthic killing, using the peroxidative oxidation and hydrogen peroxide (H$_2$O$_2$) generated by dismutation of superoxide produced during respiratory burst. In contrast, residing in the host with high level of eosinophils, helminthic worms have evolved to attenuate eosinophil-mediated tissue inflammatory responses for their survival in hosts. Our previous study demonstrated that the expression of Acan-rps-30 was significantly down-regulated in A. cantonensis L5 worms, which reside in the cerebrospinal fluid with high level of Eosinophils. Acan-RPS-30, a homologous protein of human Fau, which plays a pro-apoptotic regulatory role, may function in protecting worms from oxidative stress.

Methods: RACE, genome Walking, bioinformatics were used to isolate and analyse the structural characterisation of Acan-RPS-30; qRT-PCR and microinjection was performed to detect the expression patterns of Acan-rps-30; feeding RNAi was used to ced-3 knock-down; microinjection was performed to construct transgenic worms; oxidative stress assay was used to determine the functions of Acan-RPS-30.

Results: Our results showed that Acan-RPS-30 consisted of 130 amino acids, and was grouped into Clade V with C. elegans in phylogenetic analysis. It was expressed ubiquitously in worms and was down-regulated in both L5 and adult A. cantonensis. Worms expressing pCe-rps30::Acan-rps-30::rfp, with the refractile “button-like” apoptotic corpses, were susceptible to oxidative stress. Apoptosis genes ced-3 and ced-4 were both up-regulated in the transgenic worms. And the phenotype susceptible to oxidative stress could be converted with ced-3 defective mutation and RNAi. rps-30−/− mutant worms were resistant to oxidative stress, with ced-3 and ced-4 were both down-regulated. And the oxidative stress resistance phenotype could be rescued and inhibited by expressing pCe-rps30::Acan-rps-30::rfp in rps-30−/− mutant worms.

Conclusion: In C. elegans worms, down-regulated RPS-30 plays a defensive role against damage due to oxidative stress for worm survival by regulating ced-3 down-regulated. And this might indicate the mechanism of A. cantonensis L5 worms, with Acan-RPS-30 down-regulated, surviving in the central nervous system of human from immune attack of Eosinophil.

Background

Angiostrongylus cantonensis is a human zoonotic pathogen that may cause eosinophilic meningitis [1]. Several different hosts are required to complete the life cycle of A. cantonensis. Human, as an abnormal host, can mainly be infected by accidental ingestion of undercooked intermediate hosts, such as Pomacea canaliculata, in which the infective third-stage larvae (iL3) resides [2]. After passage to the small intestine, iL3 will penetrate the blood-brain barrier, and then infects the central nervous system, where it will develop into the fifth-stage larvae (L5), and causes angiostrongyliasis with neurological symptoms [3–6].
Eosinophils, recruited from the circulation into the central nervous system [3], are robust producers of extracellular superoxide due to expression of high level of the enzyme complex that generates superoxide [7], contributing to tissue inflammatory responses and host defense in helminthic infections [8].

Eosinophil peroxidase (EPO), contained in the granule matrix of eosinophils, would be released [9, 10]. EPO is associated with helminthic killing, using the peroxidative oxidation and hydrogen peroxide (H$_2$O$_2$) generated by dismutation of superoxide produced during respiratory burst [11–13]. In contrast, residing in the host with high level of eosinophils, helminthic worms have evolved to attenuate eosinophil-mediated tissue inflammatory responses for their survival in hosts [8]. Therefore, *A. cantonensis* L5, residing in the cerebrospinal fluid together with eosinophils, may be resistance to the damage due to oxidative stress. In our previous study, proteomic analysis of different developmental stages using two-dimensional difference gel electrophoresis (2D-DIGE) showed that the expression level of *A. cantonensis* RPS-30 (Acan-RPS-30) was lower in L5 than in iL3 [4].

*Acan-rps-30* is a homologous gene of human *fau* [14] (EBR-MuSV associated ubiquitously expressed gene), which was originally isolated from a radiation-induced osteosarcoma [15]. *Fau* is inversely inserted as the fox sequence in FBR-MuSV [16, 17], and expression of fox enhances the transformation of FBR-MuSV, presumably by inactivating *fau* expression [18, 19]. *Fau* may play important role in inhibiting tumorigenesis, with down-regulated in both breast cancer [20] and ovarian cancer [21]. Fau is also found to regulate apoptosis in human T-cell lines and HEK293/17 cells [20]. A sequence antisense to *fau* is able to decrease apoptosis induced by dexamethasone, UV or cisplatin in W7.2c cells [19]. In the parasitic nematode *Haemonchus contortus*, RPS-30 can regulate the fourth-stage larval diapauses [22]. *Fau* encodes an ubiquitin-like protein (UBiL), fused to ribosomal protein S30 (S30) as a carboxy-terminal extension [14]. These two products are thought to result from post-translational cleavage [23]. Human Fau-UBiL has 37% amino acid sequence similarity to ubiquitin and contains the C-terminal Gly-Gly dipeptide motif that participates in isopeptide bond formation between ubiquitin and lysine of target proteins [14]. However, a lack of internal lysine residues, sites of poly ubiquitin chain formation, indicates that the biological function of UBiL is different from that of ubiquitin [23]. The identification of UBiL covalently bound to Bcl-G, a member of the Bcl-2 family of apoptosis control proteins [24], suggests a pro-apoptotic regulatory role for *fau*, mediated via Bcl-G [19, 23].

Apoptosis is closely related to oxidative stress in many cell lines, mammals and the model organism *Caenorhabditis elegans* [25, 26]. Therefore, in this study, we determined the structures and functions of Acan-RPS-30 in *A. cantonensis* L5, with the aim of investigating its role in regulating oxidative stress resistance.

**Methods**

**Propagation of *A. cantonensis* and *Caenorhabditis elegans***

*A. cantonensis* ZJ strain was maintained and propagated in Wenzhou Medical University, China by cycling through the *Pomacea canaliculata* and Sprague-Dawley (SD) rats as described previously [4].
Intermediate hosts *Pomacea canaliculata* were infected with *A. cantonensis* L1 through feeding on rat feces. L3 were collected at 20 days p.i. Infected snails were shelled and crushed. The intestines and other organs were removed and the remaining tissue was homogenized. The homogenates were filtered through a 40-mesh sieve, deposited for 5 min at 4 °C, and precipitated 2–3 times at room temperature. The sediments were removed and L3 number and viability were determined by direct observation under a light microscope. Three-week-old Sprague-Dawley (SD) rats (weight 100–120 g, grade clean, Certificate SYXK [ZHE] 2005-0061), supplied by the Laboratory Animal Center of Wenzhou Medical University were orally infected with 50 L3/rat. The rats were housed in polypropylene cages with free access to food and water, and then sacrificed by anesthesia at 25 days and 45 days p.i., respectively. The L3 worms were collected from the intermediate hosts *Pomacea canaliculata*; the L5 harvested from the brains of mice (C57BL/6J (B6), Certificate SYXK (zhe2015-0009)) (non-permissive host same as humans), which were orally infected with 30 L3/mouse; the adult worms were collected from the blood vessels of the hearts and lungs. Individuals of different sexes were separated using morphological criteria: Females are usually longer and thinner than males, and males exhibit typical copulatory bursa. L3, L5, and adults were washed three times with 0.01 mol/L PBS buffer, and stored at -80 °C. These rats were not used for any other part of the study.

*Caenorhabditis elegans* strains N2, *rps-30* (tm6034/nt1) and *ced-3* (ok2734) were maintained on Nematode Growth Media (NGM) agar plates at 15 °C as described previously [27]. Worms were fed *Escherichia coli* strain OP50 unless otherwise stated. The mutant strain *ced-3* (ok2734) was obtained from the Caenorhabditis Genetic Center (CGC) (University of Minnesota, USA). The mutant strain *rps-30* (tm6034/nt1) was originally provided by Shohei Mitani, M.D., Ph.D., Department of Physiology, Tokyo Women's Medical University School of Medicine, Japan. The gene *rps-30* is essential for survival of the worms, and if the gene is deleted, worms would be sterile. Therefore, the mutant strain tm6034/nt1 was conducted as trans-heterozygous animals using a balancer nT1, which has fluorescent marker. So, fluorescence positive animals carry nT1 but animals without nT1 are mutation homozygous.

**Isolation, purification, treatment and storage of nucleic acids**

Total genomic DNA was extracted from *A. cantonensis* ZJ strain adult worms using a small-scale genomic DNA extraction kit (Takara Biotechnology Co., Ltd, Japan). Total RNA was extracted from worms at different developmental stages employing Trizol reagent (Invitrogen, USA), followed by treatment with 2 U of DNAse I (Takara Biotechnology Co., Ltd, Japan). First strand cDNA was obtained using the M-MLV RTase cDNA Synthesis Kit (Takara Biotechnology Co., Ltd, Japan). Both DNA and RNA samples were stored at -80 °C.

**Isolation of full-length cDNA and genomic DNA encoding Acan-rps-30 from A. cantonensis**

Using two degenerate primers, rps-30DF and rps-30DR (Additional file 1: Table S1), designed based on a relatively conserved S30 domain, with reference to the *C. elegans* gene (NC_003283.11) and *Homo sapiens* gene (NC_000011.10), a portion of Acan-rps-30 was amplified by PCR from cDNA synthesized from total RNA extracted from adult worms. PCR products were cloned into the pMD18-T vector (Takara Biotechnology Co., Ltd, Japan).
Biotechnology Co., Ltd, Japan) and sequenced. Based on the available sequence, gene-specific primer pairs (Additional file 1: Table S1) were designed. Then, using 5'- and 3'- rapid amplification of cDNA ends (RACE) (Takara Biotechnology Co., Ltd, Japan), two partially overlapping cDNA fragments were obtained. Products were cloned into the pMD18-T vector and sequenced. Based on these sequences, additional primers (Additional file 1: Table S1) were designed to amplify full-length *Acan-rps-30*.

Full-length genomic DNA of *Acan-rps-30* from the ZJ strain of *A. cantonensis* was obtained by a Genome Walking kit (Takara Biotechnology Co., Ltd, Japan), using primers designed based on the acquired cDNA sequence (Additional file 1: Table S1), following the manufacturer's instructions. The third-round PCR products were cloned into a pMD18-T vector and sequenced.

**Bioinformatics analysis**

A sequence alignment between *Acan-RPS-30*, *Hs-RPS-30* (NP_001988.1) and *Ce-RPS-30* (NP_505007.1) was generated using Clustal Omega. Homology models were built by SWISS-MODEL using *H. sapiens* ribosome (PDB codes 5LKS and 2L7R) as templates. Three-dimensional structural analysis was performed using the PyMOL program. All calculations were carried out under default conditions.

The amino acid sequence inferred for *Acan-RPS-30* and 7 selected other homologues sequences were subjected to phylogenetic analyses. The phylogenetic analysis was conducted using the neighbour-joining (NJ) and maximum parsimony (MP) methods, respectively, based on the Jones-Taylor-Thornton (JTT) model [28]. Confidence limits were assessed using a bootstrap procedure with 1000 pseudo-replicates for NJ and MP trees, and other settings were obtained using the default values in MEGA v.5.0. A 50% cut-off value was implemented for the consensus tree.

**Quantitative real-time PCR (qRT-PCR) analysis**

qRT-PCR was performed to determine the abundance of *Acan-rps-30* transcripts in different developmental stages (L3, L5 female, L5 male, Adult female, Adult male) of *A. cantonensis*.

Gene expression levels were determined by RT-PCR using SYBR®Green PCR Master Mix and a 7500 Real-Time PCR System (Applied Biosystems, USA). Relative gene expression was compared with 18S ribosomal RNA gene (GenBank: AY295804) as an internal loading control. The target genes and the primers used are listed in Additional file 1: Table S1. Statistical analysis was conducted using a one-way ANOVA; P < 0.05 was set as the criterion for significance.

**RNAi feeding experiments**

To generate *ced-3* specific RNAi vectors, *ced-3* cDNAs was cloned into the L4440 vector. Plasmids were transformed into *E. coli* strain HT115. Primers used for PCR analysis are listed in Additional file 1: Table S1. RNAi plates and media were prepared according to [29]. Gravid adults of *C. elegans* were allowed to lay eggs overnight on the RNAi plates and adult worms were picked off. Empty vector-containing *E. coli* were used on separate plates as negative controls.
Transgenic worms

About 2000 bp sequence upstream of Acan-rps-30 5'-UTRs was used as putative promoter. To analyze promoter activity of Acan-rps-30, the promoter regions of Acan-rps-30 and Ce-rps30 were amplified and cloned into plasmid pPD95.77 to construct pAcan-rps-30::gfp and pCe-rps30::gfp, respectively (Fig. 1a).

To perform cross-species expression of Acan-RPS-30 in N2 strain and rps-30 (tm6034/nt1) strains, cDNA sequence was amplified and cloned into pPD95.77 using the promoter of Ce-rps30 to construct plasmid pCe-rps-30::Acan-rps-30::rfp (Fig. 1b). All primers used are listed in Additional file 1: Table S1.

Recombinant plasmids were each microinjected into the gonad of young, adult C. elegans hermaphrodites as described [2, 30], together with plasmid pRF4 containing a dominant mutant allele of rol-6 gene, each at a final concentration of 50 μg/mL in the same mixture, using pPD95.77 (pCe-rps30::gfp) and pRF4 plasmid mixture as a control. The F2 and subsequent generations with a roller phenotype were analyzed and selected to examine the expression patterns of GFP or RFP, using a fluorescent microscope (Olympus IX71). A minimum of three independent lines expressing each transgene were evaluated.

Oxidative stress assay

The oxidative stress assay was performed as described previously [31]. Briefly, adult hermaphrodites (30 worms/group) were transferred to a 96-well plate containing M9 buffer with 3mM H₂O₂. After incubation at 20°C for the specified durations, the number of dead worms was determined. Worms were scored as dead when they no longer responded with movement to light prodding of the head. Three (H₂O₂) independent experiments were performed. Statistical analysis was performed with Microsoft Excel 2010 software using an unpaired two-tailed t-test.

Results

Structural characterisation of Acan-RPS-30

The complete cDNAs of Acan-rps-30 was isolated by RACE from A. cantonensis. Acan-rps-30 cDNA was 1,209 bp in length, including an open reading frame (ORF) of 393 bp (including stop codon), a 5’-untranslated region (UTR) of 190 bp, and a 3’-UTR of 626 bp (Fig. 2a). 5’ UTR harbored the consecutive pyrimidines (TTTCTTTTC), which are commonly found at the 5’ end of eukaryotic ribosomal protein mRNAs [32], and may play a role in regulating translation [33]. The 3’ UTR contained the hexamer AATAAA (positions, 612 bp downstream of the TAA). The complete Acan-rps-30 gene, isolated by Genome Walking from genomic DNA of A. cantonensis was 2,967 bp in length, consisting of 4 exons and 3 introns (Fig. 2a).

To characterize the structure of Acan-RPS-S30, sequence alignment and structural analysis were performed. The cDNA of Acan-rps-30 encoded predicted proteins of 130 amino acids (Fig. 2b), which
contained the potential cleavage sites (Gly-Gly) of the fusion protein (ubiquitin-like; UBiL-ribosome protein S30; S30). The amino acids sequence was aligned with Ce-RPS-30 and Hs-RPS-30 (Fig. 2b). The results showed that the C-terminal S30 domains were conserved (Acan-RPS-S30 versus Ce-RPS-S30 and Hs-RPS-30, 87.9% and 77.6% similarity, respectively), whereas the N-terminal UBiL domains were divergent (37.5% and 30.4% similarity, respectively). The S30 domain contained a nuclear location signal (NLS), KQEKKKKKK, with which RPS-30 can go into the nucleus and involves itself in the small subunit assembly of ribosome. Structural analysis from homology models revealed that UBiL region possessed 3 β-sheets and 2 α-helixes (Fig. 2c), and S30 region contained 2 α-helixes (Fig. 2d). The UBiL region did not harbor the K48 and K63 residues, sites of poly ubiquitin chain formation, consisted with the orthologues from other species, indicated the different functions [23], though the structure of UBiL was similar to that of ubiquitin.

**Evolutionary relationship of Acan-RPS-30 with RPS-30 orthologues from other nematode species**

To determine the evolutionary relationship between A. cantonensis and other nematodes, the predicted amino acid sequence of Acan-RPS-30 was aligned with orthologs from other nematodes, and subjected to phylogenetic analyses (Fig. 3). Acan-RPS-30 clustered closely with Dv-RPS-30 from Dictyocaulus viviparus with similarity of 89.2%. Cladistic analysis showed that the RPS-30 homologues selected from seven parasitic nematodes were mainly grouped into two clades. H. contortus, Necator americanus, D. viviparus and A. cantonensis were in Clade V; Wuchereria bancrofti, Brugia malayi and Loa loa were in Clade III. This result was in agreement with modern phylogenetic analysis of nematodes [34]. When sequences from the S30 regions only were analysed, bootstrapping did not support the clusters (data not shown). This might indicate that the divergences of the UBiL regions are likely related to species specificity.

**The expression patterns of Acan-rps-30**

To determine the relative abundance of Acan-rps-30 transcript in different developmental stages (L3, L5 and adult) and genders [females (F) and males (M)] of the life cycle of A. cantonensis, qRT-PCR was performed with the 18S ribosomal RNA gene as an internal loading control. The results showed that Acan-rps-30 was transcribed in larval and adult developmental stages examined in different levels (Fig. 4 and Additional file 2: Table S2). The expressions of Acan-rps-30 were significantly down-regulated in both L5 and adult, compared with that in L3; furthermore, the expression level in L5 was greatly lower than that in adult. This might indicate the important roles of Acan-RPS-30 in different developmental stages (L3, L5 and adult), which reside in different host.

For the lack of functional genetic and in vitro culture methods, it is unable to detect the functions of Acan-RPS-30 directly in A. cantonensis. Here, C. elegans, proposed by numerous authors as a general model for many aspects of basic molecular, cellular and developmental biology in the less tractable parasitic nematodes [34–36], was used to investigate the anatomical expression patterns of Acan-rps-30 for the closed evolutionary relationship between A. cantonensis and C. elegans, both belonging to Clade V according to Cladistic analysis [34]. Wild type C. elegans (N2 strain) were transformed with the construct
pAcan-rps-30::gfp and pCe-rps30::gfp, respectively (Fig. 1a). Plasmid pRF4 was included in all transformations as a behavioural marker. Transgenic worms showing the roller phenotype were selected. The results showed that GFP under the promoter pAcan-rps-30 was only expressed in intestine of C. elegans, mainly in the anterior end (Fig. 5a–c), which is the major tissue for lifespan regulation in C. elegans [37], in contrast to the situation in worms expressing pCe-rps-30::gfp, where GFP was expressed in almost all cells, including intestine, nervous system, pharynx, muscle (Fig. 5d–f). The different activity of pAcan-rps-30 and pCe-rps-30 might be due to the heterologous expression, with the low promoter sequences similarity (data not shown). Therefore, pCe-rps-30 was used as the promoter in following research on the functions of Acan-RPS-30 in C. elegans.

Cross-species expressions of Acan-RPS-30 in N2 and the rps-30 deletion mutant worms

In order to clarify the role of Acan-RPS-30, cross-species expression of Acan-rps-30 in C. elegans was performed. The expressing constructs containing Acan-rps-30::rfp coding sequences driven by Ce-rps-30 promoters (Fig. 1b), were used to transform C. elegans N2 strain and rps-30 deletion mutant strain (tm6034), respectively. In N2 worms, transformed with pCe-rps30::Acan-rps-30::rfp, RFP was expressed widely (Fig. 6b, c), consistent with the pCe-rps30::gfp expression pattern (Fig. 5d–f). In addition, RFP mainly focused on the nucleus for the existence of NLS in S30 region. The “button-like” morphology, corpses arising from developmental apoptosis and the gold standard for quantication of apoptosis in C. elegans [25], was seen in the anterior pharynx (Fig. 6a, d). This might suggest the pro-apoptotic effect of Acan-RPS-30, consistent with the pro-apoptotic regulatory role of Hs-RPS-30 [19, 23].

In trans-heterozygous worms (tm6034), the GFP fluorescence positive animals (pharynx), carrying nT1 are heterozygous rps-30+/− (Fig. 6e–g), and animals without GFP (nT1) are mutation homozygous rps-30−/− (Fig. 6h). After transformation of pCe-rps30::Acan-rps-30::rfp in rps-30+/− worms, the offspring contained rps-30+/− expressing pCe-rps30::Acan-rps-30::rfp (Fig. 6i–l) and rps-30−/− expressing pCe-rps30::Acan-rps-30::rfp (Fig. 6m–p), with the rps-30−/− expressing pCe-rps30::gfp (Fig. 6q–s) as the control in the following assay.

Functional role of Acan-RPS-30 in oxidative stress

To investigate the role of Acan-RPS-30 in regulating oxidative stress resistance, we performed oxidative stress assays using H$_2$O$_2$. We found that the incidence of rapid death among the N2 worms expressing pCe-rps30::Acan-rps-30::rfp was significantly higher than that among the N2 worms expressing pCe-rps30::gfp; and the rps-30 deletion mutants (rps-30−/−) was significantly more resistant than N2 worms; and this oxidative stress resistance phenotype could be rescued and inhibited by expressing pCe-rps30::Acan-rps-30::rfp in rps-30−/− mutant worms (Fig. 7a and Additional file 2: Table S2 ). This might suggest the regulating role of Acan-RPS-30 in promoting susceptibility to oxidative stress.

As oxidative stress is considered to be one of the major factors that promote apoptosis [26], we next detect the expression levels of apoptosis genes in C. elegans. The results showed that all the apoptosis
genes were down-regulated in rps-30−/− mutant worms, except akt-1 up-regulated (Fig. 7b), which inhibits CEP-1 and decreases DNA damage-induced apoptosis [38]; and ced-3 and ced-4, the core apoptosis executive genes [25], were both up-regulated in N2 worms expressing pCe-rps30::Acan-rps-30::rfp; Whereas, ced-9 down-regulated (Fig. 7c), which encodes the homologous protein to the anti-apoptotic B-cell lymphoma 2 (Bcl-2) family of proteins [39]. This might indicate the role of Acan-RPS-30 in promoting apoptosis in C. elegans.

To further determine the effect of apoptosis regulated by Acan-RPS-30 on oxidative stress susceptibility, we constructed the C. elegans strain ced-3−/−(ok2734) expressing pCe-rps30::Acan-rps-30::rfp and the strain N2; pCe-rps30::Acan-rps-30::rfp with ced-3 knocked down using RNA interference. Then, the survival percentages were detected with the strains ced-3−/− expressing pCe-rps30::gfp and N2; pCe-rps30::Acan-rps-30::rfp as controls, respectively. We found that the incidence of rapid death among the worms ced-3−/− expressing pCe-rps30::Acan-rps-30::rfp was almost the same as that among the worms ced-3−/− expressing pCe-rps30::gfp; and that among the worms N2 expressing pCe-rps30::Acan-rps-30::rfp was significantly higher than that among the worms N2 expressing pCe-rps30::Acan-rps-30::rfp with ced-3 knocked down (Fig. 7d and Additional file 2: Table S2). This might suggest that the regulating role of Acan-RPS-30 in promoting susceptibility to oxidative stress was played through CED-3, which is the core executive effector in the worm cell apoptosis [25].

**Discussion**

Eosinophilic meningitis, caused by A. cantonensis L5, is mainly attributed to the Eosinophils [40], which contribute to tissue inflammatory responses in helminthic infections [8]. Eosinophils, a well-equipped immune cell recruited from the circulation into inflammatory foci [41] and directly recognizes helminth-derived immunomodulating agents, function in host defense against helminth infection [8]. The cell surface of eosinophils possess a variety of receptors for cell signaling associated with chemotaxis, adhesion, respiratory burst, degranulation, apoptosis or survival [42], all of which may be closely associated with eosinophil-mediated tissue inflammatory responses in helminth infection [8]. Eosinophils mainly contain four main granules: crystalloid granules, primary granules, small granules, and secretory vesicles [43]. Cytotoxic granular proteins, including major basic protein (MBP), eosinophil peroxidase (EPO), eosinophil cationic protein (ECP), and eosinophil-derived neurotoxin (EDN), reside in the crystalloid granules [9, 10]. The functional role of EPO is associated with helminthic killing [44]. EPO catalyzes the peroxidative oxidation of halides and thiocyanate present in the plasma together with hydrogen peroxide (H₂O₂) generated by dismutation of superoxide produced during respiratory burst [11–13]. Eosinophils, the robust producers of extracellular superoxide due to expression of high level of the enzyme complex that generates superoxide [7], produce superoxide anions in response to helminth-derived cysteine proteases [45]. In contrast, residing in the host with high level of eosinophils, helminthic worms have evolved to attenuate eosinophil-mediated tissue inflammatory responses for their survival in hosts, such as inducing apoptosis of eosinophils [46, 47] and blocking the chemotactic effects on eosinophils [48]. Here, we identified Acan-rps-30 from A. cantonensis. The expressions of Acan-rps-30 were significantly
down-regulated in both L5 and adult. Both L5 and adult, residing in mammalian, humans and rats respectively, are attacked by immune response from hosts, such as superoxide produced by eosinophils. Our results showed that Acan-RPS-30 could promote susceptibility to hydrogen peroxide, and rps-30−/− mutant worms were resistant to oxidative stress. This might indicate the regulating function of Acan-RPS-30 in attenuating eosinophil-mediated immue attack upon L5 worms in the central nervous system of human by its lower expression. Whereas, L3 worms, with higher level of Acan-rps-30, reside in intermediate hosts (e.g., Pomacea canaliculata), in which the immune system are lower than that in mammalians, and the immune attack may be weaker, or even there may be no eosinophil-mediated superoxide attack in snails. Therefore, the higher level of Acan-rps-30 in L3 worms may indicate its multifunction in different developmental stages, such as promoting the development of L3 worms with S30 region [22]. Furthermore, the expression level in L5 was significantly lower than that in adult, which possesses the thicker cuticle than L5 larva. Though adult worms parasite in the blood vessels of the hearts and lungs of rats, in which there are the senior immune system, with attacked by superoxide from eosinophil, the thick cuticle may provide protection [49], and many other proteins may be differently expressed in the cuticle, such as the homologous gene of lec-1, which plays important role against damage due to oxidative stress [4, 49].

A. cantonensis is relatively closely related to the model organism C. elegans, both belonging to clade V [2, 34], and the homologous gene of Acan-rps-30 is Ce-rps30 (C26F1.4). Here, we used C. elegans as a surrogate to explore the in vivo functions of the homologous gene, Acan-rps-30, for the lack of effective genetic manipulation in parasitic nematode. In C. elegans, apoptosis is characterized with the refractile “button-like” morphology, apoptotic corpses, which are caused by inefficient engulfment from healthy neighboring cells [50–52]. The “button-like” appearance under differential interference contrast (DIC) optics is the gold standard for quantification of apoptosis in C. elegans [25]. In this study, the “button-like” corpses were seen in the anterior pharynx of the transgenic worm expressing pCe-rps30::Acan-rps-30::rfp, indicating the happening of apoptosis. CED-1 and CED-5 proteins can recognize corpses and are critical to engulfment [50]. The down-regulated expression of ced-1 and ced-5 in the transgenic worm expressing pCe-rps30::Acan-rps-30::rfp might contribute to the formation of corpses.

Four genes, comprising the core apoptosis pathway in C. elegans, have been identified [38, 53]. egl-1 encodes a proapoptotic BH3-only protein that antagonizes the CED-9 protein [54]. ced-9, that functions upstream of ced-4 to prevent activation of the CED-3 caspase, encodes the homologous protein to the anti-apoptotic B-cell lymphoma 2 (Bcl-2) family of proteins [39]. ced-3, encodes a proteolytic caspase protein that is activated by CED-4, the worm homolog of mammalian apoptotic protease activation factor 1 [55]. Therefore, CED-3 is the core executioner [25]. In the worms expressing pCe-rps30::Acan-rps-30::rfp, the ced-3 was up-regulated, and the worms exhibited apoptosis and susceptibility to oxidative stress; whereas, in the rps-30−/− mutant worms, the ced-3 was down-regulated, and the worms exhibited resistance to oxidative stress. This phenotype could be converted with ced-3 defective mutation and RNAi. Therefore, the function of Acan-RPS-30 in promoting susceptibility to oxidative stress might be
conducted through apoptosis by regulating CED-3. And in *A. cantonensis* L5, Acan-RPS-30 was down-regulated to enhance the resistance to oxidative stress from eosinophils for worms’ survival in host.

**Conclusions**

This study investigated the structural and functional characterisation of Acan-RPS-30 from *A. cantonensis*. We found that Acan-RPS-30 could promote worms to be susceptible to oxidative stress through apoptosis by regulating CED-3; and worms, with Acan-RPS-30 down-regulated, were resistant to oxidative stress. Our findings may reveal the mechanism for *A. cantonensis* L5 worms surviving in the central nervous system of human from immune attack of Eosinophil.

**Abbreviations**

FBR-MuSV: Finkel-Biskis-Reilly murine sarcoma virus; Gly: glycine; p.i: post infection; PBS: phosphate-buffered saline; PDB: protein data bank; GFP: green fluorescent protein; RPF: red fluorescent protein; bp: base pair; CEP-1: C. Elegans P-53-like protein; CED: CEll Death abnormality; HEK: Human Embryonic Kidney; UV: ultraviolet; Fox: FBR osteosarcoma X; ZJ: Zhejiang; AKT: Protein Kinase B; EGL: EGg Laying defective; RT-PCR, real-time PCR; RNAi, RNA interference.

**Declarations**

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**Ethics approval and consent to participate**

All the experimental animals used were treated strictly in accordance with the recommendations in the Guide for the Regulation for the Administration of Affairs Concerning Experimental Animal of the People's Republic of China. The protocol employed was approved by Laboratory Animal Ethics Committee of Wenzhou Medical College & Laboratory Animal Centre of Wenzhou Medical College (Permit Numbers: SYXK (zhe2015-0009) and SYXK [ZHE] 2005-0061). The care and maintenance of animals followed this institution's guidelines.

**Consent for publication**

Not applicable.
Availability of data and materials

Data supporting the conclusions of this article are included within the article and its additional files. The datasets used in the present study are available from the corresponding author upon reasonable request.

Competing interests

The authors declare that they have no conflicts of interests.

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Authors’ contributions

BLY, HCH and HFS conceived and designed the experiments. WWS, XMY and BLY wrote the manuscript. WWS, XMY, QS and YJZ performed the experiments. JTH and YJZ collected and analyzed the data. HCH, BLY and HFS participated in technological guidance and coordination. All authors read and approved the final manuscript.

Author details

1Department of Parasitology, School of Basic Medical Sciences, Wenzhou Medical University, Wenzhou 325035, Zhejiang, PR China. 2Department of Biochemistry, School of Basic Medical Sciences, Wenzhou Medical University, Wenzhou 325035, Zhejiang, PR China. 3Department of Pediatric Gastroenterology, The Second Affiliated Hospital and Yuying Children’s Hospital of Wenzhou Medical University, Wenzhou 325000, Zhejiang, China. 4School of First Clinic Medicine, Wenzhou Medical University, Wenzhou 325035, Zhejiang, PR China. 5Henan Provincial Engineering Laboratory of Insects Bio-reactor, China-UK-NYNU-RRes Joint Laboratory of Insect Biology, Nanyang Normal University, Nanyang 473061, PR China

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