Expression, tissue localization and serodiagnostic potential of Taenia multiceps acidic ribosomal protein P2

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Abstract

Background: The larval stage of Taenia multiceps, also known as coenurus, is the causative agent of coenurosis, which results in severe health problems in sheep, goats, cattle and other animals that negatively impact on animal husbandry. There is no reliable method to identify coenurus infected goats in the early period of infection.

Methods: We identified a full-length cDNA that encodes acidic ribosomal protein P2 from the transcriptome of T. multiceps (TmP2). Following cloning, sequencing and structural analyses were performed using bioinformatics tools. Recombinant TmP2 (rTmP2) was prokaryotically expressed and then used to test immunoreactivity and immunogenicity in immunoblotting assays. The native proteins in adult stage and coenurus were located via immunofluorescence assays, while the potential of rTmP2 for indirect ELISA-based serodiagnostics was assessed using native goat sera. In addition, 20 goats were randomly divided into a drug treatment group and a control group. Each goat was orally given mature, viable T. multiceps eggs. The drug treatment group was given 10 % praziquantel by intramuscular injection 45 days post-infection (p.i.), and all goats were screened for anti-TmP2 antibodies with the indirect ELISA method established here, once a week for 17 weeks p.i.

Results: The open reading frame (366 bp) of the target gene encodes a 12.62 kDa protein, which showed high homology to that from Taenia solium (93 % identity) and lacked a signal peptide. Immunofluorescence staining showed that TmP2 was highly localized to the parenchymatous zone of both the adult parasite and the coenurus; besides, it was widely distributed in cystic wall of coenurus. Building on good immunogenic properties, rTmP2-based ELISA exhibited a sensitivity of 95.0 % (19/20) and a specificity of 96.3 % (26/27) in detecting anti-P2 antibodies in the sera of naturally infected goats and sheep. In goats experimentally infected with T. multiceps, anti-TmP2 antibody was detectable in the control group from 3 to 10 weeks and 15 to 17 weeks p.i. In the drug-treated group, the anti-TmP2 antibody dropped below the cut-off value about 2 weeks after treatment with praziquantel and remained below this critical value until the end of the experiment.

Conclusion: The indirect ELISA method developed in this study has the potential for detection of T. multiceps infections in hosts.

Keywords: Taenia multiceps, Acidic ribosomal protein P2, Immunofluorescence, Indirect ELISA
Background

*Taenia multiceps* is a widespread parasite in many areas of the world. The larval stage, known as coenurus, mainly parasitizes the brain or spinal cord of domestic ruminants such as buffalo, cattle, goats, sheep, and yak, as well as wild species, causing lethal neurological symptoms [1–4]. Adult *T. multiceps* inhabit the small intestine of dogs, wolves, foxes and other canids [1–4]. *T. multiceps* induced coenurosis occurs almost all over the world [5], causing enormous economic losses to the livestock industry and threatening human health [1–5].

The rapid diagnosis of coenurus infection in hosts is crucial to control coenurosis and reduce its negative impact on animal husbandry. However, because *T. multiceps* infections in goats do not cause obvious early clinical symptoms, it is a significant challenge to diagnose the disease in the early stage. In recent decades, as molecular biological understanding of parasites has increased, many researchers have screened recombinant antigens for diagnosis of diseases caused by the family Taeniidae (which includes many tapeworms of medical and veterinary importance). However, data are limited on recombinant diagnostic antigens for coenurus [6–13]. Although different methods such as enzyme-linked immunosorbent assay (ELISA) [14], dot immunogold filtration assay (DIGFA) [15] and Dot-ELISA [16] have been developed for the diagnosis of coenurosis, these assays use natural worm extracts as antigens and therefore cannot be produced as commercial products. When compared with ELISA based on natural worm antigens, indirect ELISA using recombinant proteins as the capture antigen has many advantages, including high reproducibility and a stable antigen source.

Acidic ribosomal proteins are so named because of their acidic isoelectric point (pH = 3–5) and their origin in the prokaryotic or eukaryotic ribosomal large subunit. They play important roles in the maintenance of stability and activity of the ribosome [17–20], by interacting with the elongation factor involved in the regulation of protein synthesis [21–23]. Moreover, several studies have confirmed that the acidic ribosomal proteins of eukaryotes play a role in apoptosis [24, 25], the occurrence and invasion of tumors [26–28] and immune diseases [29, 30]. Acidic ribosomal proteins of eukaryotic cells are divided into three types, P0, P1 and P2 (collectively called P-proteins). The P-proteins form the lateral stalk complex of the large ribosomal subunit, comprising a 32 to 35 kDa P0 protein at the core, to which two heterodimers of acidic ribosomal proteins P1 and P2 (about 12–14 kDa each) bind, ultimately forming the stalk P0-(P1/P2)2 complex [31]. Martinez-Azorin F et al. (2008) [32] research showed that P1/P2 proteins in human cells modulate cytoplasmic translation by influencing the interaction between subunits, thereby regulating the rate of cell proliferation the recombinant P2 proteins.

The aim of this study was to characterize the *Tm*P2 protein of *T. multiceps*, determine its tissue distribution and further develop an indirect ELISA assay for the serodiagnosis of coenurosis by using recombinant *Tm*P2.

Methods

Ethics statement

All animals were raised strictly according to the animal protection laws of the People’s Republic of China (a draft of an animal protection law released on September 18, 2009). All procedures were carried out in strict accordance with the Guide for the Care and Use of Laboratory Animals by the Animal Ethics Committee of Sichuan Agricultural University (Ya’an, China) (Approval No. 2012–030).

Animals

Two healthy New Zealand white rabbits were obtained from the Laboratory Animal Center of Sichuan Agricultural University. Twenty healthy goats were obtained from a goat farm in Sichuan Province, China. All animals were fed in a barrier environment and provided with food and clean water *ad libitum*. Animals were adapted to the new environment for 1 week before the experiment.

Parasites

Adult *T. multiceps* derived from artificially infected dogs were provided by the Department of Parasitology, College of Veterinary Medicine, Sichuan Agricultural University. Coenuri were isolated from the brain of naturally infected goats in a goat farm in Sichuan Province. All materials were stored in liquid nitrogen until use.

Cloning, expression and purification of recombinant *Tm*P2

The total RNA of coenurus was extracted using a commercial kit (Huashun, Shanghai, China) and cDNA was transcribed using a RevertAid™ First Strand cDNA Synthesis Kit (MBI Fermentas, Germany) according to the manufacturers’ protocols and stored at −70 °C. Based on the transcriptome data of *T. multiceps* and the P2 sequence of *T. solium* (GenBank: L39653), gene-specific primers for *Tm*P2 were designed as follows: F1 5’-CCGGAATTCTAGCCGCTATCTCGCTGCTTAT-3’ and R1 5’-CGGCTGAGTTAGTCAAAGAGACTGAAACCCAT-3’ (Invitrogen, Carlsbad, USA), which incorporated EcoRI and XhoI restriction sites, respectively. The PCR products were digested and gel-purified (Novagen, Madison, Germany). The cDNA was subcloned into the bacterial expression vector pET-32a(+) (Novagen) and used to transform *Escherichia coli* BL21 (DE3) cells (Novagen). *E. coli* cells were cultivated at 37 °C, and then induced by Isopropyl – β – D-thiogalactopyranoside (IPTG) at an ultimate concentration...
of 1 mM. The purity of the expressed protein was measured as previously described [33].

Sequence analysis
The presence of a signal peptide was assessed using SignalP 4.1 at the Center for Biological Sequence Analysis website (http://www.cbs.dtu.dk/services/SignalP/), and cellular localization was predicted using TMHMM (http://www.cbs.dtu.dk/services/TMHMM/). The molecular weight and pI values of the predicted protein were calculated using Compute pI/Mw at ExPasy (http://web.expasy.org/protparam/).

Sera
Positive sera against parasites coenurosis (20 samples) and Cysticercus tenuicollis (7 samples) were isolated from naturally infected goats from a goat farm in Sichuan Province, and Echinococcus granulosus (8 samples) isolated from naturally infected sheep. Negative sera (24 samples) were collected from 24 cestode-free goats by autopsy. All sera were stored at −20 °C until use.

Western blot analysis
Protein extracts were prepared by homogenizing coenurosis in an NP-40 cell lysis buffer (Boster, Wuhan, China). Purified rTmP2 proteins and total worm extract were separated by SDS-PAGE and transferred onto Polyvinylidene Fluoride membranes (Boster) for 30 min in an electrophoretic transfer cell (Bio-Rad, USA). The membrane was blocked with 5 % skim milk in Tris Buffered Saline with Tween-20 (TBST) for 2 h at room temperature. Membranes were then incubated overnight at 4 °C with goat antiserum from naturally infected goats (diluted 1:100 (v/v) in 1 % skim milk in TBST). And the rest of the program was performed as described previously [34].

Immunofluorescence
To perform immunolocalization studies, T. multiceps sections were probed with specific rabbit anti-TmP2 antibodies (1:300) followed by fluorescein isothiocyanate (FITC)-conjugated goat anti-rabbit IgG (1:200; Boster, Wuhan, China) as described elsewhere [34]. The stained samples were mounted in glycerol/phosphate buffer (v/v, 9:1) and viewed under an Olympus BX50 fluorescence microscope (Olympus, Japan). Negative controls were prepared using uninfected goat serum instead of specific antibodies.

Development of the indirect ELISA
The optimal concentration of ELISA reagents (TmP2 protein and serum) was determined through standard checkerboard titration procedures [35]. Briefly, ELISAs were performed in polystyrene 96-well microtiter plates (Invitrogen) using 100 μL reaction mixtures with TmP2 protein, coated at six different concentrations (0.06, 0.12, 0.24, 0.48, 0.96, and 1.92 μg/mL) in 0.1 M carbonate buffer (pH 9.6) and incubated overnight at 4 °C. After washing three times with 0.01 M PBST, the plate was blocked with 100 μl/well of 5 % skimmed milk (skimmed milk in PBS) for 2 h at 37 °C. A serial twofold dilutions (100 μL; ranging from 1:5, 1:10, 1:20, 1:40, 1:80) of the positive and negative sera samples were used in the following step, and positive sera and negative sera were diluted in PBS. Then, after washing three times, 100 μL of HRP-labeled rabbit anti-goat IgG diluted 1:2000 in 0.01 M PBST were added to each well, followed by a 1 h incubation at 37 °C and washing three times. Finally, 100 μL tetramethylbenzidine were added into every well, and incubated at 37 °C for 20 min in the dark. After the reaction was stopped, we determined the absorbance at 450 nm in an automatic ELISA plate reader. The conditions which gave the highest P/N value and an OD450 value for positive serum close to 1.0 were defined as the optimal working conditions [36].

Determination of the cut-off value for the indirect ELISA
Twenty-four samples of coenurosis negative sera were used to assess the cut-off value under the optimal conditions for indirect ELISA. The cut-off value was calculated as the mean OD450 plus three standard deviations (SD) and will be used as a standard to identify sera positive and negative for coenurosis.
Fig. 2 (See legend on next page.)
Sensitivity and specificity analysis of indirect ELISA
The percentage sensitivity was calculated as indirect ELISA positive x 100/true positive, and the percentage specificity was calculated as indirect ELISA negative x 100/true negative. The specificity was evaluated by cross-reaction with antibody derived from E. granulosus-positive sheep and C. tenuicollis-positive goats.

Detection of anti-TmP2 antibody in goats infected with T. multiceps by ELISA
Twenty healthy adult goats were randomized to a drug treatment group and a control group (10 in each group). Each goat was orally given an average of 5500 mature, viable T. multiceps eggs. At 45 post infection (p.i.), the drug treatment group was given 10% (w/v) praziquantel by intramuscular injection at a dose of 70 mg/kg of body weight, once each day. Serum samples were collected from all the goats at weekly intervals until 17 weeks p.i.

Results
Sequence analysis, expression and reactivity of rTmP2
The TmP2 cDNA sequence consisted of an open reading frame of 366 bp encoding a putative protein with 121 amino acid residues. The protein was predicted to have a molecular weight of 12.62 kDa, a pI of 5.01, and weak hydropathicity. No signal peptides or transmembrane domains were predicted in this protein. The protein sequence of TmP2 was found to be highly homologous to those from T. solium (93% identity), Echinococcus granulosus (81%) and Hymenolepis microstoma (65%), and it exhibited homology to the acidic ribosomal protein P2 from other parasites such as Spirometra erinaceaeuropaei, Barentsia elongata and Caenorhabditis briggsae (51%, 49% and 47% identity, respectively). Recombinant TmP2 was expressed as soluble protein with a molecular weight of approximately 32 kDa (Fig. 1). Due to an additional 20-kDa epitope tag fusion peptide, the molecular mass of TmP2 was ~12 kDa, similar to that predicted from its amino acid sequence. In Western blot analysis, a positive band of 32 kDa was observed when using goat anti-T. multiceps serum, suggesting a strong reactivity of this recombinant protein (Fig. 1). No signal was present when rTmP2 was incubated with native sera (Fig. 1). In addition, the total worm extract was blotted with anti-rTmP2 rabbit serum of approximately 12 kDa (Fig. 1).

Immunolocalization of P2 protein in T. multiceps
Protein P2 was highly localized to the parenchymatous zone (PZ) of both the adult parasite and the coenurus; furthermore, it was widely distributed in cystic wall of coenurus (Fig. 2). No fluorescence staining was detected with negative sera.

Establishment of indirect ELISA
The suitability of the recombinant TmP2 protein as a diagnostic antigen was tested based on indirect ELISA. Checkerboard titration tests indicated that the conditions that gave a highest P/N value (3.64) were when the coating antigen was 0.24 μg/well and the serum dilution was 1:10 (Table 1). In the optimized test conditions, a total of 24 serum samples were analyzed to obtain the cut-off value for the indirect ELISA, and OD450 value was 0.312 with a SD of 0.0622 (data not shown). All experiments were performed in triplicate. Thus, the cut-off value was 0.499 (mean ± 3 SD). Serum samples with an absorbance ≥0.499 were scored as coenurus antibody positive,
otherwise they were determined to be coenurus antibody negative.

**Sensitivity and specificity analysis of indirect ELISA**

Specific IgG antibodies were determined in serum samples from goats infected with coenurus and *C. tenuicolli*, and from sheep infected with *E. granulosus*. (Fig. 3). Based on the cut-off of 0.499, a total of 20 serum samples from goats infected with coenurus were detected as positive, corresponding to a sensitivity of 95.0 % (19/20). There was cross-reactivity with serum from one *C. tenuicolli*-positive goat (*n* = 7) and no reactions with sera from *E. granulosus*-positive sheep (*n* = 8) and healthy goats (*n* = 12). Thus, the specificity of the ELISA using recombinant *Tm*P2 antigen to detect anti-P2 antibody was 96.3 % (26/27).

![Fig. 3 The sensitivity, specificity and cross-reactivity of indirect ELISA. The bold horizontal line indicates the cut-off value (0.499)](image)

**Fig. 4 The regularities of serological antibody against *Tm*P2 after the artificial infection of goats by *T. multiceps*. The bold horizontal line indicates the cut-off value (0.499)**
Anti-TmP2 antibody in goats artificially infected with *T. multiceps*

The regularities of serological antibodies are shown in Fig. 4 after the artificial infection of goats with *T. multiceps*. The following trends were observed: 3 weeks p.i., the drug treatment group and the control group were both positive for serum antibody to TmP2 (OD value >0.499, the cut-off value). At around 9 weeks p.i. (about 2 weeks post-injection of praziquantel), the antibody value for the drug-treated group dropped below the cut-off value; the antibody values remained below this critical value until the end of the experiment. In the control group, the antibody value dropped below the cut-off value 11 weeks p.i., but the value rose again and was detection positive from 15 weeks p.i. until the end of the experiment (17 weeks p.i.).

**Discussion**

In recent years, studies concerning acidic ribosomal P2 proteins of parasites have mainly focused on *Trypanosoma cruzi* [37–39], *Cryptosporidium parvum* [29, 40], *Toxoplasma gondii* and *Plasmodium falciparum* [41, 42]. These studies confirmed that P2 protein could induce hosts to produce a strong humoral immune response, and the protein appears to constitute a potential target for host cell invasion inhibition in both *T. gondii* and *P. falciparum* infections [29, 41, 42]. Knowledge of P2 protein is very limited for the family Taeniidae, although there are few preliminary studies on *T. solium* [43–45] which confirmed that P2 is a main pathogenic factor of human cystercerosis, and demonstrated that a P2 fusion protein expressed in *E. coli* could be used as a diagnostic antigen for human neurocystercerosis [43]. In addition, Luo et al. (2003) [44] and Su et al. (2003) [45] showed that the recombinant P2 proteins of *T. solium* expressed in *E. coli* and *Pichia pastoris* were good immunogens. In the present study, we have cloned and expressed TmP2 from coenurus for the first time.

Immunofluorescence staining showed that, in *P. falciparum*, ribosomal protein P2 (PfP2) was present at the infected erythrocyte surface at the onset of cell division, and on the merozoite surface during the *P. falciparum* host infection process [41]. In the process of *T. gondii* host cell infection, P2 protein was confirmed to be localized at the surface and in the cytoplasm of the parasite [41]. In the present study, we observed that TmP2 was highly localized to the parenchymatous zone of both the adult parasite and the coenurus; besides, it was widely distributed in cystic wall of coenurus, commensurate with its ribosomal role. Whether P2 is present in the body of *T. multiceps* oncospheres during the oncosphere host-infection process, i.e. whether the P2 protein gathers at the body surface as in *T. gondii* and *P. falciparum*, requires further study.

Due to the fact that *T. multiceps* infected goats will not show obvious clinical symptoms, it is difficult to detect in the early infective stage. This stage involves the adherence and migration of oncospheres in the major blood vessels of the intestine, and then, frequently, oncosphere transport to the nervous system including the brain and spinal cord via the circulatory system [46]. Detection of an antibody against coenurosis could be useful for early detection and treatment of the infection. ELISA, DIGFA and Dot-ELISA diagnostic methods have been established for coenurosis [14–16]. However, due to the cross-reaction of natural antibodies, previous studies have failed to establish a reliable diagnostic method. Although indirect ELISA based on a recombinant antigen against coenurosis has been developed for diagnosis, it has deficiencies including low sensitivity [47]. In our study, indirect ELISA of recombinant TmP2 was successfully established and optimized to detect coenurus in goats. The method had high sensitivity (95%) and specificity (96.3%) for 47 tested serum samples when compared with the results of necropsy. Moreover, there’s no cross reaction when using P2 protein for the detection of specificity of *E. granulosus*-positive sera. However, the indirect ELISA method against another protein (TmGST) of *T. multiceps* established based on the same sera showed one cross reaction (1/8) (unpublished). Serodiagnosis through indirect ELISA was successfully used in experimental coenurosis infection in sheep [48], however, seropositivity was observed only from the 35th day p.i. In our experiment, the antibodies could be detected by indirect ELISA in the early stage of infection (3 weeks p.i.), up to 17 weeks p.i. The anti-TmP2 serum of goats fell below the cut-off value about 2 weeks post-injection of praziquantel in the drug treatment group, and the antibody values remained below the critical value until the end of the experiment. Therefore, we conclude that indirect ELISA can be applied to the evaluation of coenurosis after the drug treatment. We did not detect the anti-TmP2 antibody between 11 and 14 weeks p.i. in the control group; however, this phenomenon was not observed when we tested antibodies against another protein (TmGST) of *T. multiceps* by indirect ELISA (data not shown). So, one can speculate that, in this period, due to decreased P2 expression, we cannot positively detect the anti-TmP2 antibody. Although this is a potential weakness of the method, it can also be applied in clinic, because infected goats have shown obvious clinical symptoms between 11 and 14 weeks p.i.

**Conclusions**

Recombinant TmP2 is a suitable diagnostic antigen for coenurus infection. TmP2-based indirect ELISA for detection of coenurus in hosts is sensitive and specific, and detects the parasite from only 3 weeks post-infection. The method will be useful for the diagnosis of
coenurosis and for validating the effectiveness of drug treatment of infections.

Competing interests

The authors declare that they have no competing interests.

Authors’ contributions

Xing Huang participated in the design of study, wrote the manuscript and performed the statistical analysis; Yingdong Yang, Yu Wang, Xiaobin Gu, Weimin Lai and Xuerong Peng contributed to animal care and experiments; Lin Chen contributed to study design and analyzed the data; Guangyoun Yang conceived of the study, participated in its design and coordination; All authors read and approved the final manuscript.

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Trypanosoma cruzi P2β protein: implications in cross-reactivity. PLoS Negl Trop Dis. 2011;5:11–10.

39. Grippo V, Niborski LL, Gomez KA, Levin MJ. Human recombinant antibodies against Trypanosoma cruzi ribosomal P2β protein. Parasitology. 2011;138:736–47.

40. Priest JW, Kwon JP, Montgomery JM, Benn C, Moss DM, Freeman AR, et al. Cloning and characterization of the acidic ribosomal protein P2 of Cryptosporidium parvum, a new 17-kilodalton antigen. Clin Vaccine Immunol. 2010;17:954–65.

41. Das S, Basu H, Korde R, Tewari R, Sharma S. Arrest of nuclear division in Plasmodium through blockage of erythrocyte surface exposed ribosomal protein P2. PLoS Pathog. 2012;8:1–19.

42. Sudasan R, Chopra RK, Khan MA, Sharma S. Ribosomal protein P2 localizes to the parasite zosite-surface and is a target for invasion inhibitory antibodies in Toxoplasma gondii and Plasmodium falciparum. Parasitol Int. 2014;64:43–9.

43. Kalinna BH, McManus DP. Cloning and characterization of a ribosomal P protein from Taenia solium, the aetiological agent of human cysticercosis. Biochem Biophys Res Commun. 1996;219:231–7.

44. Luo XN, Cai XP, Cheng HT, Liu XT. Cloning and identification of phosphoprotein P2 of Cysticercus cellulosae and expression in E. coli. Chin J Zoonoses. 2003;19:80–4.

45. Su CX, Cai XP, Han XQ, Luo XL, Zheng YD, Dou YX. Expression of phosphoprotein P2 of Cysticercus cellulosae in Pichia pastoris and its application. Sheng Wu Gong Cheng Xue Bao. 2003;19:424–7.

46. Sharma DK, Chauhanb PPS. Coenurosis status in Afro-Asian region: A review. Small Rumin Res. 2006;64:197–202.

47. An XX, Yang GY, Wang YW, Mu J, Yang AG, Gu XB, et al. Prokaryotic Expression of Tm7 Gene of T. multiceps and Establishment of Indirect ELISA Using the Expressed Protein. Acta Veterinariae Zootechnica Sinica. 2011;42:1302–8.

48. Doganay A, Biyikoglu G, Oge H. Serodiagnosis of coenurosis by ELISA in experimentally infected lambs. Acta Parasitol Turc. 1999;23:185–9.