Molecular characterization and gene expression modulation of the alternative oxidase in a scuticociliate parasite by hypoxia and mitochondrial respiration inhibitors

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Philasterides dicentrarchi is a marine benthic microaerophilic scuticociliate and an opportunistic endoparasite that can infect and cause high mortalities in cultured turbot (Scophthalmus maximus). In addition to a cytochrome pathway (CP), the ciliate can use a cyanide-insensitive respiratory pathway, which indicates the existence of an alternative oxidase (AOX) in the mitochondrion. Although AOX activity has been described in P. dicentrarchi, based on functional assay results, genetic evidence of the presence of AOX in the ciliate has not previously been reported. In this study, we conducted genomic and transcriptomic analysis of the ciliate and identified the AOX gene and its corresponding mRNA. The AOX gene (size 1,106 bp) contains four exons and three introns that generate an open reading frame of 915 bp and a protein with a predicted molecular weight of 35.6 kDa. The amino acid (aa) sequence of the AOX includes an import signal peptide targeting the mitochondria and the protein is associated with the inner membrane of the mitochondria. Bioinformatic analysis predicted that the peptide is a homodimeric glycoprotein, although monomeric forms may also appear under native conditions, with EXXH motifs associated with the diiron active centers. The aa sequences of the AOX of different P. dicentrarchi isolates are highly conserved and phylogenetically closely related to AOXs of other ciliate species, especially scuticociliates. AOX expression increased significantly during infection in the host and after the addition of CP inhibitors. This confirms the important physiological roles of AOX in respiration under conditions of low levels of O_2 and in protecting against oxidative stress generated during infection in the host.

Alternative oxidase (AOX), also known as ubiquinol-oxidoreductase, is included within the diiron carboxylate superfamily. The enzyme is characterized by cyanide resistance and is associated with maintenance of metabolic homeostasis via mitochondrial electron transport in plants subjected to various stress conditions. In plants, AOX is considered a ubiquitous enzyme; however, in algae, fungi, protozoa and several invertebrate species, it is quite sporadic. AOX is involved in the mitochondrial respiratory chain and provides an alternative pathway for the direct transfer of electrons from ubiquinone to oxygen, thus forming water. It facilitates mitochondrial respiratory activity and therefore the functioning of the Krebs cycle after inhibition of the cytochrome route due to the presence of nitric oxide, cyanide (both inhibitors of the complex IV) or antimycin A (complex III inhibitor). However, AOX reduces the energy efficiency of respiration as only 1/3 of the ATP is produced during electron transfer.
by pumping protons through complex 1\(^{17}\). In plants, AOX participates in important biological processes, such as survival and adaptation to stress factors in biotic and abiotic environments, resistance to oxidative stress and maintenance of mitochondrial and cellular energy homeostasis\(^{16,17}\). In parasitic protists, it is particularly associated with metabolic adaptation to the host and pathogenicity\(^{10,12}\). AOX is present in the mitochondria of protists such as the amoebozoans *Acanthamoeba castellani*\(^{16,22}\), *Pseudocohnilembus persalinus*\(^{26}\), *Trypanosoma brucei*\(^{26}\), *T. congolensi* and *T. evansi*\(^{27}\), *Plasmodiumfalciparum*\(^{28}\), *Cryptosporidium parvum*\(^{29}\) and *Blastoeytis hominis*\(^{16,22}\). AOX has also been detected in the Euglenozoa *Euglena gracilis*\(^{30}\), in the ciliates *Paramecium tetraurelia*\(^{31}\), *Tetrahymena pyriformis*\(^{32}\) and *Icthyophthirius multifiliis*\(^{33}\); and in the scuticociliate parasites *Pseudocohnilembus persalinus*\(^{34}\) and *Philasterides diencephali*\(^{13}\). In previous studies, we have demonstrated the presence of a cyanide-resistant alternative respiratory pathway (AP) in mitochondria of the scuticociliate parasite *P. dicensranchi*, in addition to the conventional cytochrome pathway (CP)\(^{33}\). This provided physiological evidence for the presence of a cyanide-insensitive oxidase, which was recognised by antibodies raised against a synthetic peptide derived from a fully conserved C-terminal consensus motif from plant AOX isoforms\(^{13}\). In addition, AOX inhibitors such as salicyldihydroxamic acid (SHAM) and 3,4,5-trihydroxybenzoic acid (propyl gallate, PG) significantly inhibit in vitro growth of the ciliate\(^{15,35,36}\).

Although there is clear physiological evidence for the existence of an AOX in *P. dicensranchi*, this enzyme has not previously been identified or characterized in detail at the molecular level. In the present study, we characterized the *P. dicensranchi* AOX gene and detected the AOX protein in the mitochondria. We also investigated the transcriptional modulation of this enzyme in vitro, in response to experimentally generated hypoxia, as well as in response to the presence of inhibitors of CP and AP mitochondrial respiration.

**Materials and methods**

**Experimental animals and ethics statement.** Turbot (*Scophthalmus maximus*). Juvenile fish (50 g) were obtained from a fish farm in Galicia (NW Spain). The fish were held in 200 L fibreglass tanks with a water recirculation system, at 16 °C with constant aeration and a photoperiod of 12L:12D, and were fed daily with commercial pellets (Skretting, Burgos, Spain). The fish were acclimatized to the aquarium conditions for 2 weeks before the start of the experiments.

Mice. Female CD-1 mice (Charles River Laboratories, USA) of weight 25 g were supplied by the Central Animal Husbandry Unit of the University of Santiago de Compostela (Spain).

All animal experiments and protocols were conducted in accordance with Spanish and European Legislation (R.D. 53/2013 and Council Directive 2010/63/EU) and approved by the Institutional Animal Care and Use Committee of the University of Santiago de Compostela (Spain). When required during all experiments, the animals were anaesthetized with isoflurane (mice) or 100 mg/L of tricaine methane sulphonate-MSS-222-(turbot) and finally euthanized by decapitation (mice) or by overdose of anaesthesia (turbot).

**Cell culture and experimental infections.** The isolate of *P. dicensranchi*\(^{27}\) used (II) was obtained from ascites of experimentally infected turbot (*Scophthalmus maximus*).\(^{38}\) The isolate was grown axenically at 21 °C in complete L-15 medium\(^{39}\) under (a) normoxic conditions, in culture flasks provided with vented caps to allow aeration of the culture medium, or (b) under hypoxic conditions, by bubbling the culture flasks for 10 min with argon gas and tightly closing the flask with non-vented caps, after addition or removal of the ciliates\(^{15}\).

**Sequencing of the genome and transcriptome.** For analysis of the *P. dicensranchi* genome and transcriptome, trophonts (10⁷) were concentrated by centrifugation, frozen in liquid nitrogen and sent on dry ice to Future Genomic Technologies (Leiden, Netherlands). For sequencing of the complete genome of the ciliate, a combination of short reading sequencing (Illumina technology) and long reading sequencing (Nanopore technology) was used (Oxford Nanopore Technologies). For de novo assembly of the parasite genome, the data sets were combined using the TULIP program v0.4\(^{40}\). Transcript sequences from Illumina RNA-Seq data (fragments of approximately 100 bp), obtained by amplification by SBS, were assembled using Trinity software v2.6.5\(^{41}\), included within the Galaxy application (https://usegalaxy.org/). The assembled sequences were analyzed by homology, with Blastgo 5.0 software (Biobam, Spain), and annotated. The sequences that encode proteins that are potentially related to the ciliate AOX were then selected from the *Tetrahymena thermophila* gene and protein sequences database by using the BLASTx tool Wiki TGD (https://www.ciliate.org/blast/blast_link.cgi).

**Production of recombinant AOX (rAOX) in yeast cells.** The complete nucleotide sequence that encodes *P. dicensranchi* AOX was obtained with an open reading frame (ORF) search tool (ORF Finder; https://www.ncbi.nlm.nih.gov/orffinder/), from the annotated data obtained by analysis of a RNA-Seq experiment. The codons of the original nucleotide sequence in *P. dicensranchi* were optimized using the Integrate DNA Technologies (IDT) bioinformatics tool (https://eu.idtdna.com/CodonOpt), to produce a recombinant protein in the yeast *Klyuveromyces lactis*. The AOX gene nucleotide sequence was synthesized by Invitro GeneArt Gene Synthesis (ThermoFisher Scientific), amplified by PCR with the primers FaoxKL / RaoxKL: 5′-CGC CTC GAG AGG AAA AGA ATG CAG TCC TTC GCC AGG AAA-3′/AGA ATG CGG CCC C TTA GTG GTG ATG ATG ATG ATG ATG ATG ATG ATG CTG CCG GGA GTA AGG GTC G 3′ and cloned in the integrative expression vector PKLAC2. The *K. lactis* Protein Expression kit (New England Biolabs, UK) was used to express the recombinant protein in yeast according to the manufacturer’s instructions and the protein secretion strategy, as previously described\(^{42}\).

**Obtaining anti-rAOX immune serum.** The anti-rAOX antiserum used in the immunoassays was obtained from CD-1 mice immunized intraperitoneally (i.p.) with 200 µg of rAOX. The antiserum was purified
by immobilized metal affinity chromatography (IMAC) with Ni-Sepharose and adsorbed for 30 min at room temperature in 200 μL of a 1% chitosan hydrogel adjuvant (CH) (w/v) in PBS buffer, pH 7.0. At intervals of 15 and 30 days after the first immunization, mice were reimmunized with the same dose of rAOX and adjuvant. Seven days after the second immunization, the mice were bled by decapitation and the blood thus obtained could clot overnight at 4°C. The serum was clarified by centrifugation at 2000 x g for 10 min, mixed 1:1 (v/v) with glycerol, and stored at −20°C until use.

**Sodium dodecyl sulphate polyacrylamide gel electrophoresis (SDS-PAGE) and Western-blotting.** SDS-PAGE of the rAOX protein and a ciliate lysate (CL)—obtained from cultures maintained under normoxic or hypoxic conditions—and treated under reducing conditions (after the addition of 0.02 M dithiothreitol-DTT) or under non-reducing (without the addition of DTT), was performed on linear 12.5% polyacrylamide minigels in a Mini-Protean Tetra cell system (BioRad, USA), as previously described. Once electrophoresis was completed, the gels were stained with a solution of GelCode Blue Stain Reagent (Thermo Scientific) according to the manufacturer’s instructions.

Samples separated by electrophoresis were also analyzed by Western blot, with a slightly modified version of a previously described protocol. The samples were first incubated with serum from mice immunized with AOX (anti-rAOX serum) and then with a polyclonal peroxidase-conjugated rabbit anti-mouse antibody (Dakopatts, Denmark) at 1:800 dilution. The blots were stained by adding a chromogenic enzyme substrate solution consisting of 0.003% H₂O₂ and 0.06% 3,3′-diaminobenzidine tetrahydrochloride with 0.03% NiCl₂ (DAB/NiCl₂, Sigma, USA).

**Indirect immunofluorescence (IIF).** For AOX immunolocalization in trophonts, an IIF immunoassay was performed, as previously described. Briefly, ciliates were fixed in a solution of 4% formaldehyde in PBS, permeabilized in a solution containing 0.3% Triton X-100 in PBS and blocked with a solution of 1% BSA. The ciliates were then incubated with a mouse antiserum containing anti-rAOX antibodies diluted 1:100 in PBS and with a secondary polyclonal rabbit anti-mouse immunoglobulin conjugated with fluorescein isothiocyanate (FITC; DAKO, Denmark) and diluted 1:1,000. The ciliates were then visualized by fluorescence microscopy (Zeiss Axioplan, Germany).

**Transmission electron microscopy (TEM).** For TEM analysis we followed the technique described by Parama et al. Briefly, the cultured ciliates were fixed in 2.5% (v/v) glutaraldehyde in 0.1 M cacodylate buffer at pH 7.2, post-fixed in 1% (w/v) OsO₄, pre-stained in saturated aqueous uranyl acetate, dehydrated in acetone series and embedded in Spurr’s resin. Ultrathin sections were stained with 2.5% uranyl acetate in 50% ethanol and 0.5% of lead citrate in 4.5 mM NaOH and viewed in a Jeol JEM-1011 transmission electron microscope (Jeol, Japan) at an accelerating voltage of 100 kV.

**In vitro growth assay.** To determine the growth of the ciliates under conditions of normoxia and hypoxia, we followed the protocol described by Mallo et al. Briefly, the ciliates were cultured at an initial concentration of 5 x 10⁴ trophonts/mL in complete L-15 medium in culture flasks fitted with vented caps that allowed aeration of the culture medium (normoxic conditions) and in tightly closed bottles bubbled with argon (hypoxic conditions). To ensure hypoxic conditions, the L-15 medium was bubbled with argon for 10 min until gas saturation, to completely displace the dissolved oxygen in the medium. The ciliates were then added, and the bottles were hermetically sealed to prevent entry of oxygen. To determine growth of the cultures under conditions of normoxia and hypoxia, aliquots of 100 μL were collected daily for 6 days and the number of ciliates was quantified in a Neubauer counting chamber. After collection of each sample to quantify the ciliates, the argon bubbling was repeated in all cases under the same conditions as described above to maintain the hypoxic conditions.

**Reverse transcriptase-quantitative polymerase chain reaction (RT-qPCR).** The RT-qPCR technique was basically performed as previously described, with slight variations. Briefly, the total RNA extracted from 10⁶ trophozoites/mL of *P. dicentrarchi* cultured under hypoxic/normoxic conditions, in the presence of 1 mM SHAM and 1 mM KCN (inhibitors of AOX and cytochrome-complex IV, respectively), was isolated with a NucleoSpin RNA isolation kit (Macherey–Nagel) according to the manufacturer’s instructions. The quality, purity and concentration of RNA were quantified in a NanoDrop ND-1000 Spectrophotometer (NanoDrop Technologies, USA). Two μg of sample RNA, 1·25 μM random hexamer primers (Promega) and 200 U of Moloney murine leukemia virus reverse transcriptase (MMLV; Promega) were used for cDNA synthesis (RT). Quantitative polymerase chain reaction (qPCR) analysis was performed with AOX gene-specific primers forward/reverse pair FqAOX/RqAOX (5′-CGGTTCTG CAG GGACT CCT GTG CTT AGC-3′/5′-TCGCCACT GTG TCC TCT GCC TGT TTC-3′) and *P. dicentrarchi* elongation factor 1-alpha gene (EF-1a) (GenBank accession KP952262) forward/reverse primer pair (EF1A/REF1A: 5′-TGCT CCT CTG ACC GTG CTC TCT TTG TCT CTT GTG TCC TCT TCT CTG TTG TCC TCT TCT-3′) as housekeeping gene. The reaction mixture (10 μL) used in the qPCR contained 5 μL of PowerUp SYBR Green Master Mix (Applied Biosystems), 300 nM of the primer pair, 1 μL of cDNA and RNase–DNase-free water. The qPCR was run at 95°C for 5 min, followed by 40 cycles at 95°C for 10 s and 60°C for 30 s, ending with melting-curve analysis at 95°C for 15 s, 55°C for 15 s and 95°C for 15 s in a StepOnePlus Real Time System (Thermo Fisher Scientific). Finally, relative quantification of gene expression was determined by the 2⁻ΔΔct method with the software included in the equipment.
Bioinformatic and statistical analysis. Functional analysis of proteins and classification into different families to predict the domains and important sites was carried out with INTERPRO software\(^50\). The transmembrane topology and location of signal peptide cleavage sites in amino acid (aa) sequences were predicted with the PHOBIUS program\(^51\). The N-terminal region of the protein that may contain a mitochondrial signal sequence and its cleavage site were analyzed with the MITOPROT II-v1.1 program\(^52\). The physicochemical parameters were predicted for a given protein with the The PROTPARAM tool\(^53\). The O-ß-GlcNAc attachment sites were predicted with the YINGOYANG\(^54\) and OGLCNACSCAN\(^55\) programs. For visualization of proteoforms and integration of the protein topology, the PROTTER bioinformatics tool was used\(^56\). Sub-cellular localization was predicted with the LOCTREE3 program\(^57\). The SWISS-MODEL protein server was used for modeling protein structures and complexes\(^58\). The aa sequences were aligned using the CLUSTAL OMEGA multiple sequence alignment program\(^59\), and the phylogenetic signal/noise ratio was improved with the TRIMAI tool for automated alignment trimming\(^60\). The phylogenetic trees were constructed by the Maximum Likelihood (ML) method, with the JTT model\(^61\). Branch support was given with 1,000 bootstrap replicates\(^62\) in MEGAX software\(^63\). The Bayesian inference (BI) analysis was performed with MRBAYES 3.2.6\(^64\).

The values shown in the text and figures are means ± SEM. One-way analysis of variance (ANOVA) was used for comparison of more than two samples, and the Tukey-Kramer test was used for pairwise comparisons. The Student’s t-test was used for comparison of two samples. In all cases, differences were considered significant at \(P < 0.05\).

Results

Overall structure of AOX in \textit{P. dicentrarchi}. The complete sequences of the AOX gene and cDNA were deposited in the NCBI GenBank database, with accession numbers MN193569 and MH427340, respectively. The gene encoding the AOX has a total length of 1,107 bp and contains 4 exons and 3 introns, with an average intron size of around 60 bp (Fig. 1A). The coding sequence contains 918 bp that encodes a protein of 305 aa with an estimated molecular mass of 35,638.68 daltons and a theoretical pl of 9.03. Bioinformatic analysis also indicated that this protein lacks a signal peptide; however, the MitoPro II program predicted the existence of an import signal peptide targeting the mitochondria MQSEARKFCTSSALV with a probability of 0.9503 and a cleavage site in aa 16 (Fig. 1B), as also confirmed by the bioinformatic prediction of cellular localization in the
mitochondria through the LocTree program (Fig. 1C). Bioinformatic analysis with the InterPro program (which provides functional analysis of proteins by classifying them in families and predicts important domains and sites in the sequence) showed that the protein belongs to the AOX family, i.e. that it has 6 diiron binding motifs and is O-glycosylated in the aa S3, T10, S11, T55, S117, S118, S121, T215, T259, S302 (Fig. 1B). Prediction of the biological function from the amino acid sequence of the AOX to establish the gene ontology (GO) terms indicates that this protein is associated with oxidation–reduction processes and with a biological function involving alternative oxidase activity. Analysis of the topology of the aa sequence using the Phobius bioinformatic program revealed that this protein has two transmembrane regions associated with the mitochondrial inner membrane between aa 125 and 146 and 188–211, a region located between aa 1–125 and between aa 212–305 that is located inside the mitochondrial matrix, and a region located between aa 147 and 187 that is located towards the intermembrane space of the mitochondria. However, the program for predicting transmembrane helices in proteins (TMHMM-ver. 2.0) only identified a single transmembrane region associated with the mitochondrial inner membrane between aa 188–211, a region between aa 1–188 that is located inside the mitochondrial matrix, and a region located between aa 212–305 directed towards the mitochondrial intermembrane space.

Structure of diiron active site of AOX. The protein structure modelling of AOX generated by the Swiss-Model and Swiss-PBD Viewer programs showed that the protein is homodimeric (Fig. 2A) and that each monomer contains an active site formed by two diiron centers and a OH ion (Fe$_3^+$–OH–Fe$_3^+$) (Fig. 2A,B). The computational ligand modeling indicated that Fe.1 has three ligand contacts with aa, two with glutamate (E133, E172) and one with histidine H175) residues, and one with OH; while Fe.2 interacts with three glutamate residues (E172, E223 and E279) and with OH group. Similarly, the active site of the other monomer also contains two diiron centers that interact with an OH group and with the same glutamate and histidine residues located in that monomer (Fig. 2C).

Phylogenetic analysis. We initially aligned the AOX amino acid sequences of three representative P. dicentrarchi strains (I1, B1 and C1) in order to estimate the degree of identity (Supplementary information). The AOX sequences in these P. dicentrarchi strains are highly conserved: there is a 100% identity between the AOX sequences of strains I1 and C1, and 99.67% identity with strain B1, which corresponds to a single change in a phenylalanine for a valine in aa 8 (Supplementary information). Analysis of the phylogenetic relationships between the aa sequence of the P. dicentrarchi AOX and those of AOXs of other Ciliophora indicate that are closely related to the AOX of the scuticociliate species Pseudocohnilembus persalinus (Fig. 3). The AOXs of Ciliophora species were also found to be more closely related to the AOX of Metazoan and Fungi than to Viridiplantae and other SAR species such as, for example, the Apicomplexa (Fig. 3).

Biochemical analysis and subcellular localization of P. dicentrarchi AOX. The biochemical properties of the AOX protein of P. dicentrarchi were investigated by SDS-PAGE and immunological techniques. To perform the biochemical analysis, a recombinant AOX was generated and expressed in the yeast K. lactis. The complete cDNA encoding the AOX was chemically synthesized and cloned in the yeast to produce the recombi-
nant rAOX protein, which was purified and analyzed by SDS-PAGE, verifying the size at about 35 kDa (Fig. 4B, lanes 1 and 2). Once it was confirmed that the protein is expressed in yeast, a mouse polyclonal antibody was generated for use in immunological analysis. Western blot analysis with the anti-rAOX antibody verified the existence of a single recognition band on the recombinant protein (Fig. 4A) under both reducing (lane 4) and non-reducing conditions (lane 3). However, addition to the antibody of a complete lysate of the ciliate separated under reducing conditions (Fig. 4A, lane 2) or non-reducing conditions (Fig. 4A, lane 1) led to the presence of a single 35 kDa band, in the first case, and of two bands of 35 kDa (poorly stained) and 70 kDa (intensely stained), in the second case. This indicates that the enzyme, may appear in a monomeric or dimeric state (in its native state), in the latter case, of size 75 kDa. To demonstrate the cellular location of the AOX, we performed an indirect immunofluorescence assay using the polyclonal anti-rAOX antibody (Fig. 4C). This antibody generates intense labeling immediately below the plasma membrane at the location of the mitochondrion (Fig. 4D).

Effect of hypoxia and mitochondrial respiration inhibitors on AOX expression. We first monitored the growth kinetics of *P. dicentrarchi* exposed to normoxia and hypoxia conditions. Growth of the ciliate was not altered under either condition (Fig. 5A), clearly indicating that the ciliate can grow normally under hypoxic conditions. We then analyzed AOX gene expression during different phases of growth under normoxia conditions, observing that the levels of transcripts produced under these circumstances were maintained at constant levels during the first 4 days of ciliate growth (i.e. during the logarithmic phase of growth). However, when ciliate growth reached the stationary phase, the levels AOX transcript amount increased significantly (Fig. 5B). The same was observed when the AOX protein expression was analyzed by Western blot with the polyclonal anti-rAOX antibody (Fig. 5C).
AOX gene expression during an infection in turbot was also analyzed. Turbot were experimentally infected with ciliates from normoxic cultures in the logarithmic phase of growth, and expression of the AOX gene was determined in the ciliates obtained from fish peritoneal cavity at 24 h post-infection (Fig. 5D). AOX expression was higher in ciliates obtained from infected fish than in ciliates maintained in vitro under normoxia conditions (Fig. 5D).

Finally, we carried out an experiment in which expression of the AOX gene was analyzed in ciliates maintained under normoxia and hypoxia and treated with cyanide and SHAM, classical inhibitors of mitochondrial CP and AP, respectively. The AOX transcript amount under hypoxia were significantly higher than under normoxia (Fig. 5E). Similarly, in presence of SHAM, AOX gene expression was also higher in hypoxic conditions (Fig. 5E). However, in presence of cyanide, the AOX expression levels were similar under both normoxia and hypoxia (Fig. 5E). The gene expression results were similar to those obtained by Western blot, after analysis of the AOX protein levels in the ciliates by using a polyclonal anti-rAOX antibody (Fig. 5F).

Discussion
AOX is a terminal ubiquinol oxidase that participates in an alternative mitochondrial pathway that couples oxidation of ubiquinol to the reduction of O2 to H2O without contributing to the formation of an electrochemical gradient. The first evidence for the existence of an AOX was based exclusively on physiological studies in thermogenic plant tissues, in which mitochondrial respiration resistant to inhibitors of the classical cytochrome pathway was attributed to the presence of cyanide-insensitive terminal oxidase. The subsequent use of mitochondrial purification techniques enabled isolation and purification of the AOX protein and generation of mono- and polyclonal antibodies, which led to the identification of AOX by immunochemical techniques. In recent years, molecular techniques have enabled identification of the genes that encode AOX and confirmation of the existence of this enzyme in numerous organisms. These techniques have also enabled study of the domains involved in enzymatic catalysis and of the topography of the mitochondrial membrane enzyme and its post-translational regulation, as well as reconstruction of the evolution of the enzyme in different taxonomic groups. The genes encoding AOX have been found in all higher plants and in some algae, fungi, protozoa and invertebrates, but not in vertebrates. We have found evidence of the existence of an alternative oxidase pathway in the mitochondria of the fish parasite .
and splicing, the AOX gene in *P. dicentrarchi* generated a mature mRNA that encodes a protein of an estimated molecular weight of 35,638.68 daltons, a size like the standard sizes of the AOXs of various organisms ranging between 32–36 kDa. AOX is an integral interfacial membrane protein that interacts with a single leaflet of the lipid bilayer. The first studies of the membrane localization of AOX in plant mitochondria showed that AOX has two transmembrane helices and a short loop located in the matrix. However, later topological studies indicated that AOX is an integral monotopic membrane protein that embeds into a single face (inner) of the membrane. There is some controversy regarding prediction of the transmembrane topology of the AOX in *P. dicentrarchi*. According to the Phobius program, AOX has two transmembrane helix regions, while other bioinformatic programs predict a single transmembrane helix region. In the first case, it is assumed that the C- and N-terminal domains of the protein will be located in the mitochondrial matrix; however, in the second model the N-terminal motif would be located outside the inner membrane and the C-terminal towards the mitochondrial matrix. To resolve these differences, new computational methods are probably needed, as suggested by other authors. In any case, until the crystal structure of AOX of *P. dicentrarchi* is available, it will not be possible to definitively clarify the topology of this enzyme.

**Figure 5.** Analysis of the expression of the *P. dicentrarchi* AOX gene. (A) Ciliate growth kinetics under conditions of normoxia and hypoxia. (B) AOX gene transcription levels in trophonts maintained under normoxia for 6 days. (C) Western blot analysis using a polyclonal anti-rAOX antibody and ciliated lysates obtained on different days from a normoxic culture. (D) mRNA levels in trophonts obtained from in vitro cultures under normoxia (control) conditions, and in the same ciliates after intraperitoneal injection in turbot and subsequent extraction at 24 h post-infection (24 h). (E) AOX transcript amount in trophonts maintained under normoxia and hypoxia for 6 days (Control, C) and treated with 1 mM of KCN or salicylhydroxamic acid (SHAM). (F) Western blot analysis with a polyclonal anti-rAOX antibody in ciliate lysate of trophonts maintained in the same conditions as in (E). The results presented in the graphs correspond to the mean values ± the standard error (SE; n = 5). Statistically significant difference between groups are indicated by different letters (P < 0.05).
Analysis of AOX crystal structure indicates that it is a homodimer with a non-haem diiron carboxylate protein in which the metal atoms are ligated to four fully conserved glutamate and histidine residues\textsuperscript{22,79–81}. Although the AOX family contains highly conserved active sites, small differences have been observed in ligands associated with the active center in several AOXs\textsuperscript{22}. \textit{P. dicentrarchi} has the typical C-terminal E-X-X-H motif\textsuperscript{12,92}, in which the conserved glutamate and histidine residues participate in the coordination of two iron atoms\textsuperscript{21}.

Phylogenetic studies of AOX sequences indicate that Viridiplantae, Metazoa, Fungi and SAR form monophyletic clades\textsuperscript{83}. On the other hand, Maximum Likelihood methods have revealed the existence of two main clades of AOX proteins: a clade containing AOXs of Ascomycota and Basidiomycota fungi, invertebrates, kinetoplastids and alveolate protoza, and another clade that includes plants, algae and various fungal groups\textsuperscript{43}. In our study, phylogenetic analysis by ML and BI methods also indicated the existence of two main clades of AOXs: and Sar (apicomplexans and kinetoplastids), and another including Metazoa, Fungi and Sar (Ciliophora). The present results also indicate that the AOX of \textit{P. dicentrarchi} is closely related to AOXs of other ciliate species, especially the scuticociliate \textit{P. persalinus}, but it is relatively distant from other Sar members as apicomplexans and kinetoplastids. In addition, although the enzyme sequence is highly conserved among strains of \textit{P. dicentrarchi}, some differences could be used, together with sequences of other proteins, to carry out multigenic genotyping, to enable accurate, intraspecific characterization of the isolates of this scuticociliate\textsuperscript{84}.

Plant studies indicate that AOX is an integral homodimeric membrane protein in which the monomers have an estimated molecular mass of 40 kDa and a mitochondrial directed sequence that is cleaved to generate mature forms with a molecular mass of over 32 kDa\textsuperscript{85}. Immunochemical and immunohistochemical analysis indicates that \textit{P. dicentrarchi} have monomeric forms (of about 35 kDa) and homodimeric forms (of 70 kDa) associated with the mitochondrial inner membrane.

These results strongly suggest that a disulfide bond can link two monomers (there are some Cys residues that could be involved, eg. Cys77 of AOX sequence). This is what occurs in plants, where the status of this disulfide bond impacts in enzyme activity, thus when there is not disulfide bond between two monomers (reduced form), the AOX activity is strongly activated by allosteric effectors, while the oxidized form (disulfide bond between two monomers) has little activity, even with high concentrations of the allosteric effectors\textsuperscript{79,85}. Therefore, the results obtained indicate that \textit{P. dicentrarchi} may show two AOX forms: one monomeric of 35 kDa and other dimeric of 70 kDa, due to the presence of a mixture of “reduced” and “oxidized” forms in the mitochondrial membrane.

Phylogenetic analysis of AOX from different organisms has indicated the existence of a monophyletic clade among benthic marine animals and the presence of AOX in these species, which is probably related to a mechanism of adaptation to variations in oxygen concentration, hypoxic conditions and increased hydrogen sulphide concentration\textsuperscript{86–88}. This hypothesis can also be extended to \textit{P. dicentrarchi}, a benthic (like most scuticociliates) marine organism, which is also microaerophilic\textsuperscript{13,89}.

Expression of AOX has been reported to be modulated in response to hypoxia and anoxia or after reoxygenation\textsuperscript{15}. AOX has a lower affinity for O\textsubscript{2} than cytochrome oxidase, possibly indicating higher expression/activity under conditions of low oxygen concentration\textsuperscript{89,90}. It has also been suggested that AOX expression can be increased in plants under hypoxia to reduce the formation of superoxide (O\textsubscript{2}\textsuperscript{-}), nitric oxide (NO) and peroxynitrite (ONOO\textsuperscript{-}), as well as to reduce the toxic effect of H\textsubscript{2}O\textsubscript{2}\textsuperscript{92–94}. \textit{P. dicentrarchi} is a microaerophilic microorganism capable of surviving in conditions of low oxygen concentrations\textsuperscript{12}. Under these conditions, amount of AOX transcript and of protein increase significantly in the ciliate. Gene transcription and protein expression increase during the stationary phase of growth, during which O\textsubscript{2} levels in the culture are low\textsuperscript{13}. The low concentrations of O\textsubscript{2} and the increased levels of reactive oxygen species (ROS) are also associated with infection in the host. In the first case, O\textsubscript{2} levels during the endoparasitic phase are low, while ROS levels are high as a result of the innate immune response produced during infection\textsuperscript{95,96}. Both the infection and the innate humoral immune response generated by the host induce a significant increase in AOX transcription in the ciliate.

Agents that inhibit the cytochrome bc1 significantly induce AOX transcription\textsuperscript{97,98}. Hypoxia, generated either by the removal of O\textsubscript{2} from the culture medium or by the addition of cyanide, significantly induces the transcription and expression of AOX\textsuperscript{13}. On the other hand, the addition of the AOX inhibitor, SHAM\textsuperscript{99}, only increases the transcription of the enzyme under hypoxic conditions. The data of the increase in AOX transcription obtained in this study are closely related to the results obtained at the physiological level. Thus, the determination of the mitochondrial oxygen consumption shows that under hypoxia conditions and in the presence of inhibitors of the cytochrome pathway, the ciliate preferably uses the alternative pathway respiration\textsuperscript{13}.

In conclusion, we have identified the gene and the mRNA that encode a protein belonging to the AOX family, not previously characterized at the molecular level in the scuticociliate \textit{P. dicentrarchi}. In this ciliate, the AOX occurs in the monomeric and dimeric forms and is associated with the inner membrane of the mitochondria. The amino acid sequence was found to be highly conserved among the different strains of this ciliate and is closely related to AOXs of ciliate species and, particularly, to the AOX of scuticociliates. Gene transcription increased significantly under conditions of hypoxia and afer infection and addition of inhibitors of cytochrome pathway respiration. The molecular characterization of AOX, together with the functional evidence of its role in adapting to parasitism and in resistance to oxidative stress induced after infection, opens a promising route for developing structure-based antiparasitic drug design strategies through molecular docking.

**Data availability**

The datasets generated during and/or analysed during the current study are available from the corresponding author on reasonable request.

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**Author contributions**

I.F. collected data, analysed data and revised the manuscript. J.L. designed the research and wrote and revised the manuscript. J.M.L. designed the research and wrote and revised the manuscript.

**Competing interests**

The authors declare no competing interests.

**Additional information**

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