Functional insights into the infective larval stage of \textit{Anisakis simplex} s.s., \textit{Anisakis pegreffii} and their hybrids based on gene expression patterns

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Abstract

\textbf{Background:} \textit{Anisakis simplex} sensu stricto and \textit{Anisakis pegreffii} are sibling species of nematodes parasitic on marine mammals. Zoonotic human infection with third stage infective larvae causes anisakiasis, a debilitating and potentially fatal disease. These 2 species show evidence of hybridisation in geographical areas where they are sympatric. How the species and their hybrids differ is still poorly understood.

\textbf{Results:} Third stage larvae of \textit{Anisakis simplex} s.s., \textit{Anisakis pegreffii} and hybrids were sampled from \textit{Merluccius merluccius} (Teleosti) hosts captured in waters of the FAO 27 geographical area. Specimens of each species and hybrids were distinguished with a diagnostic genetic marker (ITS). RNA was extracted from pools of 10 individuals of each taxon. Transcriptomes were generated using Illumina RNA-Seq, and assembled de novo. A joint assembly (here called merged transcriptome) of all 3 samples was also generated. The inferred transcript sets were functionally annotated and compared globally and also on subsets of secreted proteins and putative allergen families. While intermediary metabolism appeared to be typical for nematodes in the 3 evaluated taxa, their transcriptomes present strong levels of differential expression and enrichment, mainly of transcripts related to metabolic pathways and gene ontologies associated to energy metabolism and other pathways, with significant presence of excreted/secreted proteins, most of them allergens. The allergome of the 2 species and their hybrids has also been thoroughly studied; at least 74 different allergen families were identified in the transcriptomes.

\textbf{Conclusions:} \textit{A. simplex} s.s., \textit{A. pegreffii} and their hybrids differ in gene expression patterns in the L3 stage. Strong parent-of-origin effects were observed: \textit{A. pegreffii} alleles dominate in the expression patterns of hybrids albeit the latter, and \textit{A. pegreffii} also display significant differences indicating that hybrids are intermediate biological entities among their parental species, and thus of outstanding interest in the study of speciation in nematodes. Analyses of differential expression based on genes coding for secreted proteins suggests that co-infections presents different repertoires of released protein to the host environment. Both species and their hybrids, share more allergen genes than previously thought and are likely to induce overlapping disease responses.

\textbf{Keywords:} \textit{Anisakis simplex}, \textit{A. Pegreffii}, Transcriptome, Allergens, Gene expression
Background

Anisakis is a genus of nematodes (Nematoda, Rhabditida, Ascaridomorpha: Anisakidae [1]) which parasitizes marine mammals, fishes, molluscs and crustaceans. Anisakis nematodes complete their life cycle in the stomachs of cetaceans, and less frequently pinnipeds, which become infected after devouring paratenic hosts (fishes, cephalopod molluscs or krill) harboring the infective third-stage larvae (L3) [2]. Humans can also be infected by consuming raw or undercooked fish or cephalopod meat. Although L3 of Anisakis spp larvae are unable to complete their life cycle in humans, after penetration of the human gastrointestinal tract they can cause anisakiasis, a condition characterized by acute gastrointestinal manifestations of epigastralgia, nausea, abdominal pain and diarrhoea [3]. Allergic reactions such as IgE-mediated hypersensitivity, angioedema, urticaria and anaphylaxis are also associated with zoonotic exposure of humans to Anisakis spp. [4]. Fish infected with L3 are treated by freezing, which kills the larvae. However, while infection by live L3 is necessary for many of the symptoms of anisakiasis, humans sensitized to Anisakis may show symptoms after consuming fish that has been correctly frozen, cooked and even processed [5–7]. Many Anisakis spp. proteins maintain their allergenic properties even after thermal treatment [7–9]. Episodes of allergic disease in the absence of recent allergen exposure have also been also described [10]. Anisakiasis has also become an important epidemiological problem. More than 20,000 cases of anisakiasis have been reported worldwide since 1960 [11], with higher incidences in areas such as Japan, the Netherlands, France, Spain, Germany and California where raw fish is traditionally or increasingly being eaten. Given the widespread consumption of raw fish in the form of sushi and sashimi, Japan has however the highest prevalence of gastric anisakiasis. In Spain, where Anisakis spp. are considered to be an emerging health problem [6], infection frequency ranks between 0.43 and 22% in fish that reach the market have been reported, with very high rates of infection in particular species [12].

The allergenic potential of Anisakis spp. resides in their allergome (or full set of encoded allergen products). Anisakis spp. contains at least 15 distinct types of major allergens (Ani s 1 to Ani s 14 and Ani s 11.0201) according to the Allergen Nomenclature site (http://www.allergen.org) [13, 14]. Up to 28 different proteins described as potential new allergens were identified by proteomic analysis of A. simplex s.s. and A. pegreffii [15]. A transcriptomic analysis reported 36 and 39 allergens in A. simplex s.s. and A. pegreffii respectively [16].

Morphological taxonomy of Anisakis is based on excretory system and esophageal intestinal region of L3 [17, 18] and the morphology of adult males [19]. As sampling of adult Anisakis from within definitive hosts is problematic, and L3 diagnosis is difficult, a molecular taxonomic approach for L3, using the nuclear ribosomal internal transcribed spacers (ITS), has been developed [20]. These markers have revealed both potential additional species diversity within Anisakis as well as recombinant genotypes [21–23]. The genus Anisakis is commonly accepted to include 10 named species: A. simplex sensu stricto (s.s.), A. berlandi (formerly A. simplex C), A. pegreffii, A. typica, A. ziphidarum, A. physeteris, A. brevispiculata, A. paggioe, A. nascettii and A. schupakovi. At least 2 additional undescribed species (Anisakis sp. 1 and Anisakis sp. 2) have also been designated from L3 genotypes [24, 25].

The described species are A. simplex s.s., A. pegreffii and A. berlandi: these constitute the Anisakis simplex sensu lato (s.l.) complex [26–28]. A. simplex s.s. and A. pegreffii have been identified as the main etiological agents of anisakiasis [29–33] but it remains unclear if these 2 species are equally pathogenic. Importantly, A. simplex s.s. and A. pegreffii are able of hybridizing in the areas where they are geographically sympatric [34, 35] and co-infect the same fish host [36]. The described “species” with recombinant genotypes may be the products of interspecific hybridization [34].

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The described “species” with recombinant genotypes may be the products of interspecific hybridization [34]. While no fertile adult hybrids have been found, L3 with genotypes suggestive of hybridisation between A. simplex s.s. and A. pegreffii have been identified [24]. These hybrids may express distinct suites of allergenic and immunoreactive proteins [15].

Here we present a transcriptomic analysis of A. simplex s.s., A. pegreffii and their hybrids sampled in the same development stage (L3) from the same host (Merluccius merluccius) captured in waters where A. simplex s.s. and A. pegreffii are sympatric. We used these data to compare their metabolic profiles, gene ontology profiles and expression patterns. We also present an online database, AnisakisDB, which provides access to the abundant data we have generated.

Methods

Taxonomic identification and selection of specimens for RNA sequencing

Larvae specimens in the same development stage (third stage larvae L3) of Anisakis were obtained from the FAO 27 area of fishing distribution into a general survey carried out by our laboratory. L3 were extracted from hosts following published procedures [34, 37, 38]. L3 were rinsed in 0.9% saline solution, placed in an antibiotic solution for 30 min and then rinsed in bi-distilled water for 1 h before molecular identification and RNA extraction. The larvae were individualized (from each host) and for each specimen the caudal part was used for DNA extraction and PCR amplification for species identification; this caudal part and the rest of the body were
separately stored at -80 °C until required. Species identification in that survey was performed following the taxonomic criteria of [23, 39] by using the ITS1 region of the nuclear ribosomal DNA (rDNA). Briefly; individual *Anisakis spp.* L3 stage juveniles was placed in an Eppendorf tube after previously having a small part of the caudal region removed to allow molecular identification of each individual. DNA was extracted and purified using the Speedtools Tissue DNA Extraction Kit (Biotools) following manufactures instructions. Molecular identification was carried out for each individual using PCR-RFLP. The forward primer A 5'-GTCGAATTCGTAGGTGAACCTGCGGAAG GATCA-3' and reverse primer B 5'-GCGGATCC GAATCTGGTATGTTTCTTTTCCCT -3' [39] were used in reactions containing, 10 mm Tris-HCl (pH 8.3), 1.5 mm MgCl2, 50 mm KCl, 200 mm each of dATP, dCTP, dGTP and dTTP and 1 unit of DNA polymerase (Biotools B and M labs, S.A. Madrid, Spain). Initial denaturation was carried out for a period of 2 min at 94 °C followed by 35 cycles of 94 °C for 1 min, 58 °C for 1 min, 72 °C for 1 min followed by a final 7 min extension at 72 °C. Amplified DNA fragments were digested with the restriction enzymes HhaI and HinFl (New England Biolabs, Massachusetts, MA, USA) following manufacturer's instructions. Restriction fragments were separated by electrophoresis in Tris- Borate-EDTA (TBE) buffered 2.5% agarose gel, stained with SYBERsafe and visualized with UV illumination.

**Specimens selection for transcriptomics**

The detected hybrids can differ as the species (*A. simplex* s.s. or *A. pegreffii*) contributing the maternal and paternal genomes, which can influence their expression patterns. We have used mitochondrial DNA sequences, which are inherited maternally, to differentiate the two types of hybrids. We obtained partial sequences of the mitochondrial cytochrome oxidase subunit 2 (COII) gene [40] from each hybrid and determined its maternal species by comparing the COII sequence with a set of COII sequences obtained from 100 populations of *Anisakis*, using molecular phylogenetic methods. The COII gene was amplified as detailed in [40]. The obtained sequences were pooled with 41 other *Anisakis* COII sequences downloaded from GenBank [41], and then aligned using the ClustalW program [42] as implemented in Bioedit [43]. Phylogenetic relationship of populations was obtained by Maximum Likelihood and Bayesian Tree (Evolution Model: GTR + I + G) [44] (Additional file 1) considering as outgroup the sequences of *Contracaecum* sp. and *Toxocara canis*. The Bayesian information criterion (BIC) as implemented in jModelTest v2.1.4 [45] selected GTR + I + G (I = 0.4950; G = 0.6230) as the evolutionary model that best fit the data. The selected model and model parameters were used in the maximum likelihood (ML) analysis performed with PhyML v.3.1 [46].

The robustness of the inferred trees was tested by nonparametric bootstraping (BP) using 1000 pseudo replicates. Bayesian inference was also performed with MrBayes v.3.2.5 [44], running for $1 \times 10^7$ generations (four simultaneous Markov chains; sample frequency 100). Four independent partitioned analyses were performed and checked for stationarity and convergence of the chains with Tracer v1.6 [47]. Three data partitions were analyzed: first, second, and third codon positions. Model parameters were estimated independently for each one of the respective data partitions. Burn-in was set to the first 1,000,000 generations. The robustness of the inferred Bayesian trees was determined using Bayesian posterior probabilities (BPP; as obtained from majority-rule consensus trees of the post burn-in trees) (provided as Additional file 1).

In order to minimize a possible host effect, the larvae used for the three RNA sequencing experiment were obtained from ten different individuals of the same fish host (European hake, *Merluccius merluccius*) and when it was possible, from fishes which were parasited at the same time by *A. simplex* s.s., *A. pegreffii* and their hybrid haplotypes (labeled by yellow and green colors in Additional file 2).

RNA samples from the 3 groups of selected specimens were pooled and 500 ng of each processed using the Illumina TruSeq RNA sample prep kit v.2 (Illumina) to generate 2 poly(A)-selected libraries, one with small inserts (SI library; size range = 200–325 bp) and one with large inserts (LI library; size range = 260–840). Paired-end reads (2 × 100 bases) were generated on Illumina HiSeq 2000. A total of 19–23 million pass-filter read pairs were obtained from each library. Raw reads were converted to fastq format using Casava on the BaseSpace platform [48]. Read set qualities were assessed using FastQC [49], and trimmed of adapters and low quality data with Cutadapt [50] and Prinseq [51]. An assembly was produced independently for each species (*A. simplex* s.s. and *A. pegreffii*) and for the hybrids. The SI and LI libraries trimmed fastq files from each specimen group were assembled de novo using Oases [52, 53] using a kmer range of 21–31 and following the specifications indicated in the Oases manual. All the assembly metrics provided in Table 1 were obtained using the Perl script Assemblathon stats [54] with the exception of the number of unigenes, which was provided by Oases.

**Sequence annotations**

The transcriptomes of *A. simplex* s.s., *A. pegreffii* and the hybrids were annotated using the NCBI tool blastx [55] with an evalue of 10e-5 as cutoff threshold and the NCBI non-redundant protein (NR) and the Eukaryotic Orthologous Groups (KOG) databases [56] as reference subjects. Gene Ontology (GO) terms as well as their evidence code [57] and Enzyme Commission (EC) numbers [58] were
assigned to annotations using correlation tables among GenBank accessions and GO and EC information. InterPro domain [59] annotations were also annotated using identical methodology. Transcripts that had no informative blastx matches were further used to search the NCBI nucleotide database (NT) using NCBI BLAST tool blastn and identical evalue cut-off. The annotation protocol is automatized as a pipeline cited in the section below “pipelines”. Proteins (open reading frames) longer than 75 residues were predicted from transcripts using Transeq from the EMBOSS package [60] and OrfPredictor [61]. Signal4.1 [62] and TMMHMM v.2.0 [63] were used to predict secretory signal peptides and transmembrane regions. Metabolic pathways were retrieved from the KEGG web site [64] using EC annotations as queries.

Reference transcriptome

Minimus2 from the Amos package [65] was used to merge the 3 assemblies into a merged transcriptome. Remaining redundancy was eliminated using CD-HIT [66] at a similarity threshold of 0.95. The merged transcriptome after redundancy filter resulted in 75,380 consensus transcripts that were annotated via BLAST as described above for transcriptomes using the NR database.

Differential expression and enrichment analyses

Bowtie2 [67] was used to map the two RNAseq pair-end libraries sequenced per each *Anisakis* sample on the merged transcriptome. In all cases over 97% of the reads were successfully mapped on the reference (data not shown). Next, Corset [68] was used to parse the bam files resulting from the mapping step and extract a cluster file grouping the 75,380 consensus transcripts into 74,751 transcript clusters and a count file summarizing the read counts obtained per each cluster from all pair-end libraries mapped to the reference merged transcriptome. The count file was used as input to EdgeR [69] to perform 3 differential expression tests at the whole-transcriptome level: “hybrids vs. *A. pegreffii*,” “hybrids vs. *A. simplex s.s.*” and “*A. pegreffii* vs. *A. simplex s.s.*”. The SI and LI pair-end libraries prepared from each sample were considered as technical replicates. Transcripts were considered as differentially expressed when they had a *p*-value < 0.05 after False Discovery Rate (FDR) correction applied by EdgeR using the Benjamini-Hochberg method [70] (referred as just FDR through the rest of the manuscript). Transcripts clusters were assigned the annotations of their consensus transcripts using the GPRO worksheet [71]. Clusters from transcripts with no NR/NT annotations were considered in differential expression analyses only if they had non-zero read counts in at least 2 of the 3 samples.

GO and metabolic pathway enrichment analyses were performed using GOSeq [72] and the results from differential expression as input, following the GOSeq manual indications for non-native transcriptomes. Concretely, we used 4 input files: 1) a file summarizing the consensus transcripts found as differentially expressed as a FDR < 0.05 as a cutoff in differential expression analyses; 2) a file with all assayed consensus transcripts; 3) a file with the GOs or with the metabolic maps associated to the consensus transcripts (depending on the analysis); and 4) a file with the sequence size of each assayed consensus transcript. A priori, enriched GO terms and metabolic pathways supported by *P*-values < 0.05 in the resulting Wallenius distribution, were considered as significant. A 5% FDR correction was performed on the *P*-values obtained from the enrichment analyses using the Benjamini-Hochberg method to outline the most enriched GOs and pathways.

### Table 1: Metrics for de novo assembly transcriptomes

| Summary | *A. simplex s.s.* | *A. pegreffii* | Hybrid | Merged |
|---------|------------------|---------------|--------|--------|
| Total transcriptome size | 88,007,524 | 68,071,234 | 50,568,936 | 67,459,080 |
| Unigenes (Loci) | 36,645 | 31,988 | 29,566 | -NA- |
| Transcripts (isoforms) | 121,907 | 91,541 | 76,848 | 75,380 |
| Longest transcript (bp) | 10,774 | 11,724 | 10,798 | 16,240 |
| shortest transcript (bp) | 100 | 100 | 100 | 107 |
| % transcripts >1Kb | 20.3% | 21.7% | 17.9% | 28.7% |
| N50 (bp) | 973 | 1026 | 885 | 1276 |
| L50 (bp) | 25,878 | 19,118 | 16,598 | 15,385 |
| %A | 30.87% | 30.25% | 30.11% | 30.41% |
| %C | 19.60% | 20.03% | 20.36% | 18.64% |
| %G | 19.64% | 19.98% | 20.49% | 20.60% |
| %T | 29.89% | 29.74% | 29.04% | 29.87% |
| %Ns | 0% | 0% | 0% | 0% |
With the aim to also assess differential expression of genes encoding for excreted or secreted (ES) compounds read counts of transcript clusters with annotation of secretory leader peptide from Signal-P domain or with annotation of GO terms for extracellular compartments were exported to a new count file. Counts in this file from transcripts sharing the same protein descriptions (from NR annotations) were combined using a custom php script called combine_counts.php that is available in AnisakisDB (see the section below AnisakisDB). We combined information from the Uniprot browser [73] and the literature to filter the transcript set to include only those verified as coding for ES products or with confirmed secretory signal peptides but unknown sub-localization. The final count file had 356 non-redundant protein secretory leader peptide from Signal-P domain or with an-EC thus creating abundant material (annotations and other material such as scripts, statistics graphical rep-ECations and the differential expression and enrichment files). To house the aforesaid material and make it freely av-ECailable for readers, we constructed a web-available data-ECbase, AnisakisDB. The database was programmed using Lar-ECavel framework for public client platform and a CodeIgniter 2 framework for server platform, both based on the PHP programming language [74], a MySQL data-ECbase [75] and an Apache server [76] hosted in a Linux en-ECvironment. The database portal includes a blast search powered by the NCBI BLAST package [77]. Within Ani-ECakisDB, sequences, annotations and backups of all performed analyses of differential expression and enrichment are distributed in seven sections (“Transcriptomes”, “Blast”, “Download”, “COII markers”, “Secretome” and Allergome) managed by an intuitive menu and additional utilities to retrieve or compare sequences and annotations. In addition, within the section “Transcriptomes” we have implemented different web site sections providing access to 5 independent Venn Diagrams created based on the GI, KOG, InterPro, GO, EC annotations used to create the Venn diagram of Fig. 1. In each Venn-based web site, clicking on any intersected number the Venn opens a dia-EClog summarizing the annotations that correspond to the intersection. The Section “COII markers” is another web site where the COII-based phylogeny inferred for selection of specimens has been graphically implemented, also as a dynamic tree representation. AnisakisDB is freely and on-ECline accessible at www.anisakis.mnecn.csic.es.

Results and discussion
Transcriptome de novo assembly and annotation
We sampled Anisakis L3 from a single ocean region (FAO 27) and fish species (M. merluccius), to avoid is-Diaissues that might arise from response to different parate-ECnic hosts and minimize other environmental and phylogeographic confounding factors. L3 were identified ac-ECcording the taxonomic approaches of [23, 39] by using the ITS1 region from rDNA. The phylogenetic tree of 141 COII sequences from 100 populations plus another 41 sequences obtained from GenBank showed all A. sim-plex s.s and all A. pegreffii samples clustering in separate sister clades (Additional file 1). The tree also showed that all COII sequences from hybrids fell into one or an-ECother of these two clades, allowing immediate classification of hybrids as to their maternal species. RNA was extracted from pools of 10 A. simplex s.s. 10 A. pegreffii and 10 hybrids (five mothered by A. simplex s.s. and five mothered by A. pegreffii). Two paired-end RNA-Seq datasets were generated from each pooled sample, forming technical replicates of the transcriptome. Four transcriptome assemblies were generated, one for each sample (A. simplex s.s., A. pegreffii and hybrids) and one
that merged the 3 sample-derived ones (Table 1). The 3 sample-derived transcriptomes are similar in span and complexity. The assembly of *A. simplex s.s.* was longer and had an elevated number of predicted transcripts, apparently because of a greater number of isoforms (30,366 more than *A. pegreffii* assembly and 45,059 more than the hybrid assembly). We think this might be due to sequencing technical issues, elevated levels of alternative splicing (i.e. isoforms) or elevated heterozygosity in the transcriptome of *A. simplex s.s.* as the number of unigenes is in contrast quite equilibrated (4657 unigenes more than *A. pegreffii* assembly and 6989 more than the hybrid assembly).

**Sequence accessions**

COII sequences have been deposited at European Nucleotide Archive (ENA) under the accession numbers LT883269 - LT883368 (Additional file 2) (For more

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**Fig. 1** Annotations. a Blast top hits per subject species using NR as refseq database for unigene annotation. b Distribution of annotated and unannotated transcripts and unigenes annotated via blast using NR as refseq database. c Venn diagram showing unique and shared features of non-redundant annotations per transcriptome (Gs, KOGs, InterPros, GOs, ECs)
Table 2 Summary of annotations per transcriptome

| Sequences with annotations | A. simplex s.s. | A. pegreffii | Hybrids |
|----------------------------|----------------|--------------|---------|
| Gene Identifiers (GIs) from NR/NT | 62,483 transcripts, 14,057 unigenes | 60,816 transcripts, 14,393 unigenes | 53,741 transcripts, 14,183 unigenes |
| KOG clusters                | 38,717 | 29,977 | 28,114 | 9928 |
| InterPro domains            | 17,093 | 13,514 | 12,884 | 5125 |
| GO Terms                    | 14,701 | 11,773 | 11,101 | 4501 |
| ECs                         | 5416 | 4237 | 3913 | 1617 |
| Predicted proteins          | 49,755 | 41,324 | 38,782 | 13,834 |
| Predicted signal-P domains  | 1124 | 603 | 482 | 318 |
| Predicted TM domains        | 11,613 | 8906 | 7545 | 4109 |

Expression patterns and enrichment of metabolic pathways and gene ontologies

The merged (consensus) transcriptome reconstructed in this study based on the three Anisakis transcriptome consist of 75,380 consensus sequences, which were used as a mapping reference to compare the expression patterns at the whole-transcriptome level of the 3 taxa. Of the 75,380 consensus transcripts, 32,999 have no annotation and therefore correspond to unknown sequences expressed by the at least two of three taxa (they might be unknown proteins, potential non-coding RNAs and/or even mobile elements). The remaining 42,381 consensus transcripts correspond to coding sequences annotated based on 12,511 different proteins (according to the number of non redundant GI annotations) thus meaning that any biological interpretation of differential expression and enrichment based on the merged transcriptome is based on the expression patterns of 12,511 coding genes (around 68% of the pan-transcriptome here characterized based on the three transcriptomes). After mapping the reference transcriptome, the 75,380
consensus sequences of the merged transcriptome were grouped into 74,751 transcript clusters (see Methods) correcting any potential bias in isoforms as detected in the assembly of A. simplex s.s. The Biological Coefficient of Variation (BCV) and the average dispersion of the assayed samples were inferred and evaluated to assess suitability of the fastq libraries, resulting in quite appropriate values (BCV = 0.4359 and dispersion = 0.19005) for a study of differential expression. A similar conclusion resulted from plotting BCV against the average log of counts per million of mapped reads. Additionally, a multidimensional scaling (MDS) plot based on logFC of the differences among taxa sources and the sequenced replicates per taxa shows that while the SI and LI libraries used as taxon replicates fall close to one another, the replicates of different taxa fall widely separated (Fig. 2b). This result anticipates important differences among species and hybrids. The first plot dimension correspond to the differences among taxa sources (A. simplex s.s., Hybrids, A. pegreffii) that provides the major source of variability. The second plot dimension corresponds to the sequenced replicates per taxa (small and medium insert size fastqs) that present practically no differences with the exception of the hybrids where the 2 replicates are slightly more heterogeneous because they were created as a pool of 5 hybrids close to A. simplex s.s., and 5 other close to A. pegreffii. Consistent with this perspective, of these 74,751 transcript clusters, 8239 resulted as differentially expressed in the analysis performed between the hybrids and A. pegreffii using FDR < 0.05 as cutoff, 23,549 were differentially expressed in the analysis performed between the hybrids and A. simplex s.s. using same FDR, and 24,813 were differentially expressed between A. pegreffii and A. simplex s.s. with same FDR (Fig. 2a and Additional file 3). In general, these analyses of differential expression based on the merged transcriptome suggest that the A. pegreffii parental expression pattern predominates over A. simplex s.s. in the hybrids, in agreement with population genetic studies of Moroccan Anisakis [36].

The 272 ECs identified as common in the 3 transcriptomes link to 53 metabolic pathways [64] related to nutritional requirements and biosynthesis, energy metabolism, metabolism of xenobiotics, environmental information processing, signal transduction and genetic information processing. Ten of these 53 pathways were shown as differentially enriched in at least one of the 3 enrichment comparisons after correction of the pvalue at FDR < 0.05 (Additional file 4). As shown in Fig. 3, two of these pathways (oxidative phosphorylation and methane metabolism) were differentially enriched in all comparisons thus indicating that the main metabolic differences among A. simplex s.s., A. pegreffii and their hybrids concern to the energy metabolism. Both A. pegreffii and A. simplex s.s. present overexpression with respect to the hybrids as for transcripts associated to methane metabolism while differentially expressed transcripts involved in oxidative phosphorylation are overexpressed in A. pegreffii with respect to the 2 other taxa. Figure 3 also reveals that the enrichment of the glycosaminoglycan degradation pathway was significant between the performed comparisons of the hybrids and A. pegreffii against A. simplex s.s. but not when the hybrids and A. pegreffii are compared. The degradation of glycosaminoglycans pathway mainly concerns to transcripts coding for hyaluronoglucosaminidases, which are known to play role in the extracellular matrix as soluble components and polyelectrolytes interacting with growth factors and other transient components of the extracellular matrix [78]. In this case, the enrichment derives from a strong overexpression of the transcripts coding for these enzymes in the hybrids and A. pegreffii with respect to A. simplex s.s. Reversely, the pathway of alanine aspartate and glutamate metabolism is shown as significantly enriched when the hybrids and A. simplex s.s. are compared with A. pegreffii but not when they were compared each to other thus suggesting that the A. pegreffii parental does not predominate in all expression patterns of hybrids. The remaining pathways (pyruvate metabolism, aminoacyl-tRNA biosynthesis, streptomycin biosynthesis, starch and sucrose metabolism, pentose phosphate pathway, and cysteine and methionine metabolism) were found as differentially enriched only in 1 of the 3 performed comparisons, predominantly when A. pegreffii was compared against any of the 2 other taxa (Fig. 3). Results suggest that A. pegreffii and the hybrids exhibit for these pathways more differentially expressed transcripts than those that would be expected by chance. In fact, the ratio between differentially expressed and assayed transcripts associated to the enrichment of each pathway is predominantly lower when the hybrids are compared with A. pegreffii than when they are compared with A. simplex s.s. or when A. simplex s.s. is compared with A. pegreffii. Finally, it is worth to also discuss the detection of streptomycin biosynthesis pathway, which is associated with the over expression of 7 transcripts encoding for 2 type of enzymes, inositol-3-phosphate synthases (lyases) and dTDP-4-dehydrorhomannose 3,5-epimerases (isomerases) in A. pegreffii and A. simplex s.s. when compared to the hybrids. The expression of the genes coding for these enzymes might have been induced by the antibiotic treatment applied on L3 specimens before the extraction (see methods) thus apparently revealing some deficiencies in the hybrids with respect to their parents of for this metabolic pathway. However, we should to also take into account that Inositol-3-phosphate synthase also participates in the metabolism of inositol phosphates, mainly to produce membrane receptor playing role in signalling, cell
growth, apoptosis, cell migration, endocytosis, and cell differentiation [79], while dTDP-4-dehydrorhamnose 3,5-epimerases are also associated to the nucleotide sugar metabolism and with the biosynthesis of dTDP-rhamnose to produce rhamnose. The biosynthesis of dTDP-rhamnose is a new pathway involved in hypodermal development or in the production of the cuticle or surface coat in larval stages of *C. elegans* and other nematodes [80]. We thus think that inositol-3-phosphate synthases and dTDP-4-dehydrorhamnose 3,5-epimerases of anisakids might link not only to the metabolism of secondary metabolites but also to signalling, nucleotide sugar metabolism and larval development.

Analyses of GO enrichment were also performed based on the 1211 GOs shared by the 3 transcriptomes and revealed differential enrichment of 91 GOs at FDR < 0.05 in at least on of the 3 performed comparisons (Additional file 4). Forty of these 91 correspond to biological processes terms, 37 are molecular functions and 14 cellular components (Fig. 4). Thirteen of these 91 were significant in all comparisons and correspond with functions, processes and sub-localizations associated with the energy metabolism and this mainly includes oxidases, oxido-reductases, dehydrogenases and other enzymes and binding proteins with assigned roles in redox processes, and electron carriers probably related to the transmembrane electrochemical gradient as well as roles for heme-biding proteins (Cytochrome P450s). We think that the latter might also be related with sterol and heme recruitment. As other nematodes, anisakids probably

![Fig. 2 Transcriptome differential expression patterns. a BCV versus the average log of counts per million (CPM). b MDS plot based on logFC of differences among taxa sources and the sequenced replicates per taxa (small and medium insert size fastqs). c MA-plots (one per each differential expression analysis) representing the log Fold Change (logFC) against the log-average log CPM per each transcript cluster across each pair of compared samples. Differentially expressed clusters supported by FDR < 0.05 are plotted in red. Each MA-plot is accompanied by a summarization of the results of each differential expression comparison.](image-url)
Recruit exogenous sterol and heme recruited from host sources to implement sterol and heme dietary needs. Consistent with this argument, the 3 transcriptomes addressed in this study present annotations of prenyltransferases, squalene oxidases and sterol regulatory element-binding proteins (SREBP); prenyltransferases play role in the synthesis of pyrophosphate and geranylpyrophosphate, a major metabolic chokepoint in de novo sterol biosynthesis pathway [81] while squalene and derivatives are the main source of host sterol and SREBP is known to stimulate sterol biosynthesis. The enrichment of the remaining GOs shown in Fig. 4 was significant in 1 or 2 of the 3 performed comparisons and refers to molecular functions biological processes and cellular components related to the enriched pathways shown and discussed above but also with transposition (mainly integrases and proteases encoded by Bel/Pao LTR retrotransposons), transport, growth and development, locomotion, and with the extracellular region (ES proteins, allergens and cuticle collagens). In broad terms and for almost all evaluated GOs (with several exceptions) the ratio between differentially expressed and assayed transcripts is again lower if the hybrids are compared with A. pegreffii than when the hybrids are compared with A. simplex s.s. or when the latter is compared with A. pegreffii. In turn, the comparison between the hybrids and A. pegreffii presents largely more enrichment than the others; of the 91 GOs shown as differentially enriched, 67 terms are significant in the aforesaid comparison between the hybrids and A. pegreffii while 27 GOs are significant when the hybrids and A. simplex s.s compared and 34 when the latter is compared with A.
Interestingly, the most prominent differences in GO enrichment observed between the hybrids and the 2 parents derive from the differential expression of transcripts related with the inositol polyphosphate 5-phosphatase activity involved in inositol phosphate-mediated signaling. While these 2 GO terms are significantly enriched when they are compared with their parental species but no when the parents are compared each to other. Indeed, we think that the deregulation of the inositol polyphosphate 5-phosphatase enzyme and how it affects the whole inositol phosphate-mediated signaling pathway is worth to be investigated, moreover if we take into account that hybrids are not able to reach adult stages and that inositol phosphate-mediated signalling has been associated to longevity, growth and reproduction in nematodes and other invertebrates [82]. The enrichment of GOs is also consistent with the perspectives provided by the differential expression analyses and the enrichment of metabolic pathways that together supports the idea of that although the alleles of the A. pegreffii parental apparently predominate over those of A. simplex s.s. in the expression patterns of L3 larval stage hybrids the latter are intermediate biological entities with metabolic and functional requirements, quantitatively different to those of the two parental species.

The set of metabolic pathways and GOs (i.e. the functional profiles) of A. simplex s.s., A. pegreffii and their
hybrids are consistent with the aerobic intermediary metabolism of nematodes thus supporting the idea that almost all pathways typically observed in *C. elegans* and other nematodes are present also in these anisakids. A graphical idealisation of the intermediary metabolism of nematodes is represented in Fig. 5 according to [81–85] and the most representative pathways and GOs detected in the 3 *Anisakis* transcriptomes have been highlighted. Our conclusion is that the *A. pegreffii* parental predominates over *A. simplex* s.s. in the expression of hybrids albeit the latter, and *A. pegreffii* also present significant differences between them indicating that hybrids are intermediate biological entities with metabolic and functional requirements quantitatively different to those of their parental. The hybrids thus offer an excellent opportunity to investigate the mechanisms of speciation in *Anisakis* including the novel biological and ecological processes where hybrid genotypes might have a relevant role or significant lacks. Hybridization among *A. simplex* s.s. and *A. pegreffii* has indeed an important role in the evolutionary biology of these 2 species through gene introgression [86]. While hybrids do not appear to reach the definitive host, allowing development to reproductive adults, they reflect the dynamics of moving areas of sympatry. Hybrids may thus be crucial in understanding the microevolutionary processes active in *A. simplex* s.l., in host-parasite ecology (including the range of paratenic hosts exploited), and in whale migrations (as interbreeding must take place in whales as final host).

**Genes encoding for excretory/secretory proteins**

To specifically focus on the potential sub-localizations of the products encoded by differentially expressed transcripts associated to enrichment we created a box plot based on the logFCs of transcripts associated to enriched cellular component GOs (Fig. 6). The analysis shows the relationships of over/under expression observed among the 3 taxa with respect to products sub-localized in mitochondrion, respiratory chain, ATP transporting factors and myosin complexes with ATP-dependent roles in muscle contraction, motility and the extracellular region. A significant number of outliers (due to their extremely high logFCs) correspond to transcripts whose product is sub-localized in the extracellular region thus indicating relevant differences with respect to genes encoding likely

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**Fig. 5** Nematode intermediary metabolism, as described for *C. elegans* and other species. The main pathways and GOs found as enriched among the 3 *Anisakis* transcriptomes are highlighted.
ES proteins. All ascaridids, *Anisakis* included, release ES products to the host environment. Infective L3 penetrate the intestinal wall of the fish host, and may encyst on the surface of the internal organs or migrate to the muscular tissues. The migration may occur not only when the host is alive [87–91] but also after death [92–94]. These processes are likely to be mediated by secreted enzymes. ES proteins have additional functional roles connected to nutrition, infectivity, allergy, immune evasion or pathogenicity [34, 95, 96] playing important roles in the host-parasite interaction. In the aforeshown analyses performed at the whole-transcriptome level we showed that part of the differential enrichment between taxa corresponds to transcripts encoding likely ES proteins (above). According to our annotations these transcripts correspond to the expression of at least 356 non-redundant gene descriptions of which we create a count file of mapped reads to explore how different are the expression patterns of the investigated taxa with respect to ES genes (see Methods). Nearly three-quarters (266/356) were found to be significantly differentially expressed in one or more comparisons under a FDR < 0.05 as threshold cutoff (Additional file 5). More specifically, 151 genes were found as differentially expressed when comparing hybrids and *A. pegreffii*. Of these, 79 genes were overexpressed in hybrids and 72 overexpressed in *A. pegreffii*. Comparing hybrids with *A. simplex s.s.*, identified 191 differentially expressed genes, of which 107 were overexpressed in hybrids and 84 in *A. simplex s.s.*. Comparing *A. simplex s.s.* and *A. pegreffii*, 100 transcripts were overexpressed in *A. pegreffii* and 82 in *A. simplex s.s.*. A heatmap with clustering of the logFC per comparison of the 70 most differentially expressed ES genes (Fig. 7) permits to visualize four major patterns of expression. Rows (i.e. the gene expression patterns) split in 4 Clusters of expression designated as clusters 1,2,3 and 4. Cluster 1 groups genes over-expressed in *A. pegreffii* compared to both *A. simplex s.s.* and hybrids, cluster 2 is a single gene, overexpressed in hybrids. Cluster 3 is defined by genes over-expressed in *A. simplex s.s.* compared to both *A. pegreffii* and hybrids, and genes belonging to Cluster 4 were predominantly over-expressed in both *A. pegreffii* and hybrids compared to *A. simplex s.s.*. ES genes with allergenic properties are highlighted with red asterisks. In particular, the analysis shows that in mixed infections different species and hybrids of *Anisakis* may produce different kinds and levels of ES proteins, or in other words that the modulation of the host environment may depend on the mix of larvae from species and hybrids. Considering that hybrids are not able to complete their life cycle and that no colony of L3 hybrids has been yet described in nature parasitizing alone their hosts (as far as we know hybrids have been only reported accompanied by their parental species) it is tempting to speculate based on our results that hybrids might be able to survive in host
environments only when they are accompanied by the parental species. Facilitation may also derive from genes overexpressed or downexpressed in the hybrids.

**Detection and classification of allergens**

Anisakiasis is a consequence of allergic reaction to several allergenic molecules expressed and/or secreted by *Anisakis* in their intermediate hosts. Briefly, 15 allergens (Ani s 1 to Ani s 14 and Ani s troponin C) are formally considered from *A. simplex s.s.* according to the AllFam database of allergen families and the WHO/IUIS Allergen Nomenclature Database [13, 14]. Five of the 15 *Anisakis* formal allergens - Ani s1, and Ani s2 and Ani s3, Ani s7, and Ani s12 - are considered to be major...
allergens (one to which >50% of allergic patients immunologically react and present unusual levels of IgE and IgG in serum [97, 98]); Ani s 1 belongs to the animal Kunitz serine protease inhibitor family of protease inhibitors (AF003); Ani s 2 is a myosin heavy chain family (AF100) [99] that includes the highly cross-reactive mite paramyosin major allergens [100]; Ani s 3 is part of the tropomyosin family (AF054), which have been identified as minor inhalative allergens in arthropods and as major food allergens in crustaceans, mollusks and Anisakis [101]; and Ani s 7 and Ani s 12 are 2 proteins with unknown biological functions. The remaining Anisakis allergens are minor allergens; Ani s 4 belongs to the cystatin family (AF005) [102] of reversibly binding cysteine protease inhibitors; Ani s 6 belongs to the cysteine-rich trypsin inhibitor-like domain family [103] (AF027); Ani s 5, Ani s 8 and Ani s 9 are members of the nematode-specific SXP/RAL-2 family (AF137), which have unknown function [104, 105]; Ani s 10, Ani s 11 and Ani s 14 similarly remain functionally unclassified [106–108]; Anis s troponin C is a member of the EF hand family (AF007), which has roles in calcium signaling and binding [109]; and Ani s 13 belongs to the globin family (AF009) [110]. Yet more interesting, recent proteomic analyses have identified up to 28 potential new allergens in Anisakis that can be classified into distinct allergen families [15] and a more recent transcriptomics study [16] has identified several new allergens from A. simplex s.s. and A. pegreffii. Additional potential allergens (Ani s 24 kDa, Ani s CCOS3, Ani s cytochrome B, Ani s FBPP, Ani s NADHDS4L, Ani s NARA5 and Ani s PEPB) have been also considered according to previous studies [111–113]. All these molecules are classified or can be classified as food allergens because they are present in marine fish. The characterization of allergens in anisakids is still open and that will aid diagnosis and possible alleviation of the serious symptoms of anisakiasis. A complete allergome (all allergen proteins) catalogue for anisakids is required.

Special importance should be paid to proteins related to mechanisms of parasitism and pathogenicity of these nematodes. According to a recent transcriptomic study [114] in the pharyngeal region of A. simplex s.s. there are at least 226 transcripts which are potentially involved in invasion and host pathogen interplay. The same study found at least 339 transcripts with a similar role in A. pegreffii. Regarding this, 13 immunoreactive proteins from the esophageal glands of A. simplex s.s. have been described as potential allergens (data not published; PARASITE project [115]).

We created a protein sequence database including 509 non-redundant food allergens from fungi, plants and animals (see Methods). This database was used to search homologues in the Anisakis consensus transcriptome we had assembled in this study. We detected high-scoring matches (E-values < 1.99E-06) to 937 consensus transcripts from 121 different allergens that can be classified into 74 families of allergens following the criteria of classification contemplated by AllFamDB and WHO/IUIS 13,14 and other classificatory criteria [111–113] (Table 3; Additional file 6). Of the 74 identified families, 41 were detected through similarity to allergens described from animals (15 correspond to all Anisakis formal allergens), 15 were detected using fungal allergens and 13 using allergens of plants (i.e. some of these had best matches with allergens from fungi and plants). The remaining 5 sequences were identified using allergens present in more than one realm of life. Differential expression analyses revealed the 74 families to be present in the 3 transcriptomes (Additional file 6). Thirty one families were found as differentially expressed (under a FDR threshold of 0.05) when comparing hybrids to A. pegreffii, 33 families when comparing hybrids to A. simplex s.s., and 36 families when comparing A. pegreffii to A. simplex s.s. These expression patterns are consistent although more slightly with the other analyses performed by us (the A. pegreffii parental genotype slightly predominates over A. simplex s.s. in the hybrids’ expression patterns). It is striking that all 3 taxa express a large number of different allergens types, many more than has been described previously. So the potential allergen repertoire of these anisakids can be expanded. We think that this repertory might be related to (or be able to partially explain) the cross-reaction of anisakiasis patients to allergens from other sources. It is usually assumed that after sensitization by nematode L3, subsequent contact with these same allergens is required to induce an allergenic response [116]. A number of Anisakis allergens are ES proteins [15], with high affinity to specific immunoglobulin E (IgE) [117], and patients primed by A. simplex s.s. might respond adversely to cross-reactive antigens of not only other anisakids but also of other nematodes and arthropods [118]. Somatic allergens such as Ani s 3 (tropomyosin) cross-react with proteins of crustaceans, mollusks and insects (particularly the American cockroach) [119].

Conclusions

A. simplex s.s., A. pegreffii and their hybrids show differences in their gene expression patterns in the larval L3 stage. Strong parent-of-origin effects were observed indicating that hybrids are intermediate biological entities among their parental species, and thus of great interest in the study of speciation in nematodes. Differential expression analyses based on genes coding for secreted proteins suggests that co-infections present different repertoires of released proteins to the host environment. Both species and their hybrids, share more allergen genes than previously thought and are likely to induce
Table 3 Distribution of Allergen Families (AllFams) and logFC results from differential expression

| Allergen Families                  | FamID | LogFCs Hh vs Ap | LogFCs Ap vs As | Con Seqs | Allergen Name | Realm | Type of allergen |
|-----------------------------------|-------|----------------|----------------|----------|---------------|-------|-----------------|
| 24 kDa protein                    | Not yet assigned | 2,24,615 | 0,94,729 | -1,29,887 | 20 | NA | Animal | NA |
| 60S acidic ribosomal protein      | AF070 | 0,70,555 | 0,25,574 | -0,44,982 | 4 | Asp f 8, Alt a 12, Asp f 8, Fuc f 1 | Fungi | Minor |
| Aldehyde dehydrogenase            | AF040 | 0,06989 | -0,10,196 | -0,17,186 | 26 | Blo t 4 | Animal | Minor |
| Alpha-amyrase                     | AF033 | -0,05472 | 0,32,169 | 0,37,641 | 6 | Alt a 10, Cla h 10 | Fungi | Minor |
| Ani s 10 allergen                 | Not yet assigned | 1,07632 | 0,06603 | -1,01029 | 1 | Ani s 10 | Animal | NA |
| Ani s 11 allergen                 | Not yet assigned | -2,67,162 | 2,26,799 | 4,93,961 | 6 | Ani s 11, Ani s 11 L1, Ani s 11 L2 | Animal | NA |
| Ani s 12 allergen                 | Not yet assigned | -0,22,028 | -0,25,132 | -0,03103 | 4 | Ani s 12 | Animal | Major |
| Animal Kunitz serine protease inhibitor | AF003 | 1,50,377 | -1,17,032 | -2,67,407 | 9 | Ani s 1 | Animal | Major |
| ARM-like                          | Not yet assigned | 0,88,344 | -4,73,273 | -5,61,615 | 44 | Ani s 7, Ani s 14 | Animal | NA |
| ATP:guanido phosphotransferase    | AF049 | 0,28,397 | 0,03604 | -0,24,793 | 5 | Pen m 2, Der p 20, Plo i 1 | Animal | Major, Minor |
| ATP synthase                      | Not yet assigned | 0,41,629 | 0,19,251 | -0,22,378 | 3 | Box d OSCP | Animal | Minor |
| BCL7 family                       | AF121 | -0,42,399 | 0,29,518 | 0,71,916 | 1 | Horn s 3 | Animal | NA |
| Calreticulin family               | AF055 | -0,27,705 | -0,37,927 | 0,65,631 | 7 | Pen ch 31 | Animal | NA |
| CAP family                        | AF044 | -0,63,714 | -1,22,008 | -0,58,293 | 12 | sol i 3, Dol a 5, Vesp c5, Poly s 5, Pol d 5, Dol m 5, allrgn_V5/Tpx1 | Animal | Major, NA |
| Carboxylesterase                  | AF140 | -0,07,113 | -0,09,074 | -0,01,961 | 50 | Api m 8 | Animal | Minor |
| Casein kinase II regulatory subunit | Not yet assigned | -0,54,905 | -0,81,674 | -0,26,768 | 2 | Gal d Phosvitin | Animal | NA |
| Catalase                          | AF047 | 1,53,912 | 0,81,466 | -0,72,447 | 1 | Pen c 30 | Fungi | Minor |
| Chitinase class III and peritrophin-like | AF077 | -0,55,642 | -0,22,104 | 0,33,539 | 12 | Der p 15, Der f 15 | Animal | Major |
| Collagen                          | AF097 | -0,20,422 | 0,10,867 | 0,31,289 | 2 | Bos d alpha 2 A | Animal | NA |
| Cyclophilin                       | AF038 | 0,18,030 | 0,27,158 | 0,09,912 | 13 | Asp f 27 | Fungi | NA |
| Cystatin                          | AF005 | 2,62,999 | -0,32,427 | -2,95,425 | 4 | Ani s 4 | Animal | NA |
| Cysteine-rich trypsin inhibitor-like domain | AF027 | -2,05,519 | 0,03,836 | 2,09,355 | 5 | Ani s 6 | Animal | NA |
| Cytochrome b-c1                    | Not yet assigned | 0,59,193 | 0,94,815 | 0,35,621 | 6 | NA | Animal | NA |
| Cytochrome B5                      | Not yet assigned | 1,95,134 | 0,37,530 | -1,57,620 | 3 | NA | Animal | NA |
| Cytochrome c                       | AF006 | 0,14,607 | 1,04,446 | 0,89,838 | 5 | Cur i 3 | Fungi | Major |
| EF hand family                     | AF007 | 0,20,495 | 0,40,417 | 0,19,922 | 34 | B1a g 6, Per a 6, Lit v 3, Ani s Troponin C | Animal | Major, Minor, NA |
| Enolase                           | AF031 | -0,52,169 | -0,22,686 | 0,29,484 | 5 | rho m 1, hev b 9, Alt a 6 | Fungi, Plant | Minor |
| Eukaryotic aspartyl protease       | AF004 | -0,11,780 | -0,35,758 | -0,23,979 | 14 | Bla g 2 | Animal | Major |
| Eukaryotic elongation factor 1     | AF011 | 0,03,983 | -0,05,788 | -0,09,771 | 8 | Pen c 24 | Fungi | NA |
| Fe/Mn superoxide dismutase         | AF020 | 0,00,817 | 0,19,412 | 0,18,595 | 2 | Hev b 10 | Plant | Minor |
| Gelsolin family                    | AF074 | -0,81,825 | -0,61,973 | 0,19,852 | 17 | Der f 16 | Animal | Minor |
| Allergen Families                | FamID  | LogFCs Hh vs Ap | LogFCs Ap vs As | Con Seqs | Allergen Name | Realm | Type of allergen |
|---------------------------------|--------|----------------|----------------|----------|---------------|-------|-----------------|
| GILT family                     | AF155  | 1.61, 144      | 0.92, 869      | −0.68, 276 | Tri a 27      | Plant | NA              |
| Globin                          | AF009  | 2.89, 321      | 0.55, 871      | −2.33, 450 | Ani s 13      | Animal| Minor           |
| Glutathione S-transferase        | AF010  | 0.06592        | 0.47, 962      | 0.41, 370 | Bla g 5, Der p 8 | Animal| Major           |
| Glyoxalase superfamily          | AF082  | −0.40, 931     | 0.77, 379      | 1.18, 310 | Ory Glyoxalase I | Plant | NA              |
| GMC oxidoreductase              | AF081  | 0.37, 676      | 0.45, 602      | 0.07, 926  | Mala s 12     | Fungi | Major           |
| Heat shock protein Hsp70        | AF002  | −0.17, 513     | −0.64, 037     | −0.46, 524 | Pen c 10, Cla h HSP70 | Fungi | Minor           |
| Heat shock protein Hsp90        | AF042  | −0.19, 335     | 0.11, 856      | 0.31, 191 | Asp f 12      | Fungi | NA              |
| Hevein-like domain              | AF043  | 0.35, 419      | −0.39, 541     | −0.74, 952 | Hev b 11      | Plant | NA              |
| Histidine acid phosphatase      | AF062  | −0.03, 153     | −0.32, 299     | −0.29, 146 | Api m 3       | Animal| Minor           |
| Hylauronidase                   | AF103  | 1.02, 016      | −0.32, 364     | −1.34, 380 | Pol a 2, Api m 2 | Animal| NA, Major       |
| Intermediate filament protein   | AF008  | −0.30, 492     | −0.89, 969     | −0.59, 477 | Horn s 5      | Animal| NA              |
| Lactate/malate dehydrogenase    | AF014  | −0.78, 551     | −0.56, 985     | 0.21, 566 | Cit la MDH    | Plant | Major           |
| Larval allergen (Brugia malayii)| Unclassified | 0.32, 808 | 1.48, 199 | 1.15, 313 | NA            | Animal| NA              |
| Lipase                          | AF037  | 0.09, 487      | −0.64, 847     | −0.74, 333 | Dol m 1, Pol d 1, rhi o Lipase, The l lipase | Animal, Plant | Major, NA |
| Lipocalin                       | AF015  | 0.51, 1183     | 0.05, 554      | −0.45, 629 | Tyr p 13      | Animal| Minor           |
| Major allergen (Loa loa)        | Unclassified | −0.00, 868 | −0.22, 329 | −0.21, 461 | NA            | Animal| NA              |
| Myosin heavy chain              | AF100  | −0.21, 310     | 0.07, 600      | 0.28, 910 | Ani s 2, Blo t 11, Der f 11, der p 11 | Animal, Major | Minor       |
| NAC domain                      | AF107  | 0.58, 475      | 0.39, 473      | −0.19, 002 | Horn s 2      | Animal| NA              |
| Nitrophorin                     | Unclassified | 0.27, 507 | 0.31, 344 | 0.03, 836 | Cim I Nitrophorin | Animal | NA              |
| Papain-like cysteine protease   | AF030  | −0.01, 475     | 0.08, 466      | 0.09, 941 | Act d 1, Ana c 2, car p 1, Gly m Bd 30 k, blo t 2 | Animal, Plant | Major, NA |
| Patatin family                  | AF104  | −0.27, 069     | −3.25, 600     | −2.97, 961 | Hev b 7       | Plant | Major           |
| Phosphoglycerate kinase         | Unclassified | −0.22, 025 | −0.55, 632 | −0.33, 067 | Can a PGK     | Fungi | NA              |
| Profilin family                 | AF051  | 0.19, 390      | −0.08, 435     | −0.27, 824 | Par j 3       | Plant | Major           |
| Prolamin superfamily            | AF050  | 0.07, 050      | −0.07, 100     | −0.14, 148 | Tri a 26      | Plant | NA              |
| Prolyl oligopeptidase family    | AF061  | −0.04, 633     | 0.30, 623      | 0.35, 255 | Ves v 3, Tri r 4 | Animal, Fungi | Major, NA |
| Proteasome subunit beta type    | Unclassified | 0.98, 838 | 1.03, 285 | 0.04, 012 | Zea m 20S     | Plant | Minor           |
| Protein kinase                  | AF017  | −0.43, 020     | −0.00, 592     | 0.42, 428 | Sal k 2       | Plant | Major           |
| Ribosomal protein L3            | AF058  | 1.01, 034      | 1.22, 333      | 0.21, 599 | Asp f 23      | Fungi | Minor           |
| SART-1 family                   | AF116  | −0.01, 191     | −0.21, 899     | −0.20, 797 | Horn s 1      | Animal| NA              |
| Serpin serine protease inhibitor| AF018  | 0.84, 445      | 0.38, 305      | −0.46, 140 | Gal d 2       | Animal| Major           |
| Short-chain dehydrogenase       | AF028  | −0.00, 468     | 0.02, 563      | 0.12, 031 | Alt a 8, Cla h 8 | Fungi | Major, Minor    |
| Subtilisin-like serine protease  | AF021  | 0.06, 703      | −1.01, 133     | −1.08, 002 | Rho m 2       | Fungi | Major           |
| SXP/RAL-2 family                | AF137  | 1.79, 413      | 0.74, 628      | −1.04, 786 | Ani s 8, Ani s 9, Ani s 5 | Animal | Minor |
| Thioredoxin                     | AF023  | 0.57, 209      | 0.74, 939      | 0.17, 728 | Asp f 28      | Fungi | Major           |
| Translationally controlled tumour protein | AF136 | 0.40, 520 | −0.40, 064 | −0.80, 584 | Horn s TCTP | Animal | NA |
| Triosephosphate isomerase       | AF032  | 0.01, 199      | 0.03, 933      | 0.02, 734 | Tri a 31      | Plant | Minor           |
overlapping disease responses. In the protein sequence database which includes 509 non-redundant food allergens from fungi, plants and animals we detected 121 different allergens belonging to 74 allergen families following recognized classificatory criteria.

Additional files

**Additional file 1:** Phylogenetic relationships of the considered populations as an individual’s selection basis for RNA sequencing of *A. simplex* s.s., *A. pegreffii* and their hybrid haplotype. Tree was obtained based on maximum likelihood (ML) and Bayesian inference analysis using mitochondrial COII gene and the GTR + I + G evolutionary model. Numbers at nodes correspond to ML bootstrap proportions (BP) (above number) and Bayesian posterior probabilities (BPP) (under number). (PNG 621 kb)

**Additional file 2:** List of species and specimen used in the phylogenetic tree of Additional file 1. Code of the voucher specimen and accession number for mitochondrial gene COII (*: sequences obtained from GenBank). Labeled are the specimens selected for RNA sequencing (first number, population; second number specimen). *A. simplex* s.s. – *A. pegreffii* refers to hybrids haplotype according Ab ello et al. [23]. (DOCX 47 kb)

**Additional file 3:** Differential expression analyses at the whole transcriptome level. Results obtained from the analyses of differential expression; 1) Hybrids vs. *A. pegreffii*, 2) Hybrids vs. *A. simplex* s.s.; 3) *A. pegreffii* vs. *A. simplex* s.s. Rows summarizing significant differentially expressed transcripts supported by FDR < 0.05 are highlighted in light blue. (XLSX 12115 kb)

**Additional file 4:** Differentially enriched metabolic pathways and GOs. Annotation of metabolic pathways and GOs integrating results of differential enrichment using G2Seq. (XLSX 232 kb)

**Additional file 5:** Differential expression of ES transcripts. 1) Count file used as input in the differential expression test performed at the ES gene level; 2) Summarization of the 356 ES genes accompanied by logFC values and their FDR support obtained from the 3 performed differential expression analyses and additional information relative to consensus sequences grouped to each gene description, as well as associated GOs terms and identification of SignalP domain. This summarization is also available in AnisakisDB as a dynamic table from where sequences and annotations can be navigated and retrieved. (XLSX 91 kb)

**Additional file 6:** Anisakis allergome. 1) Count file and results of differential expression for the 74 allergen families detected in *A. simplex* s.s., *A. pegreffii* and hybrids. 2) Blast results obtained in the search performed using 509 known allergens as queries against the reference *Anisakis* transcriptome. Nine hundred and thirty seven consensus sequences of the reference transcriptome were detected to significantly match to the queries and were classified into 74 allergen families of which we include additional information. This summarization is also available in AnisakisDB as a dynamic table from where sequences and annotations can be easily navigated and retrieved. (XLSX 351 kb)

**Abbreviations**

**AllFam**: Allergen family; **BCV**: Biological coefficient of variation; **BIC**: Bayesian information criterion; **BPP**: Bayesian posterior probability; **COII**: Cytochrome oxidase subunit II; **CPM**: Counts per million; **EC**: Enzyme commission; **ENA**: European Nucleotide Archive; **ES**: Excretory/secretory; **FDR**: False discovery rate; **G2**: Gene Expression; **GII**: Hybrid haplotype; **IgE**: Immunoglobulin E; **ITS**: Internal transcribed spacers; **KOB**: Eukaryotic orthologous groups; **L3**: Third-stage larva; **logFC**: Log2 fold change; **MDS**: Multidimensional scaling; **ML**: Maximum likelihood; **NR**: NCBI non-redundant protein database; **NT**: NCBI nucleotide database; **ORF**: Open reading frame; **TM**: Transmembrane domain

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**Availability of data and materials**

COII sequences have been deposited at European Nucleotide Archive (ENA) under the accession numbers LT883269 - LT883368 (For more details, see the section AnisakisDB in Methods). Raw read data have been deposited at the NCBI SRA Study SRP072976, BioProject PRJNA316941 and BioSamples SAMN04592605, SAMN04592630, and SAMN04592599. Assemblies, sequences, annotations and backups of all analyses performed in this study are freely available in AnisakisDB at www.anisakis.mncn.csic.es.

**Authors’ contributions**

AN, LR, SCA, MB, IM, MGM, MC, JPA and CL conceived and designed the study. AN and JPA coordinated the study. SCA, NCS, LR, RR, YJR, MC and MGM performed the biological and taxonomic experiments based on COII signal. LR, SCA, NCS and RR, performed the RNAseq experiments. CL, LR, SC,
SCA, VJR, RR and AN performed the bioinformatics analyses. RF, CL, and AN created the database. CL, AN, BS, LR, SCA, MB, IM, MGMT, MC analyzed and interpreted the data. CL, AN and MB wrote the manuscript. All authors read and approved the final manuscript.

Competing interest
The authors declare that they have no competing interests.

Ethics approval and consent to participate
Not applicable. Anisakis samples were collected during a survey of the FAO 27 area in the fish central Madrid Market. No ethical approval or field permissions were necessary.

Consent for publication
Not applicable.

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